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# CAR LIGAND-BINDING DOMAIN POLYPEPTIDE CO-CRYSTALLIZED WITH A LIGAND, AND METHODS OF DESIGNING LIGANDS THAT MODULATE CAR ACTIVITY

# 5 <u>Technical Field</u>

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The present invention relates generally to the structure of the ligand-binding domain of CAR, and more particularly to the structure of the ligand-binding domain of CAR in complex with a ligand. The present invention also relates to CAR binding compounds and to the design of compounds that bind to CAR.

# **Abbreviations**

•	amu	-	atomic mass unit(s)
15	ATP	-	adenosine triphosphate
	ADP	-	adenosine diphosphate
	BSA	-	bovine serum albumin
	CaMV	-	cauliflower mosaic virus
20	CAR	-	constitutive androstane receptor
	$CAR\alpha$	-	constitutive androstane receptor alpha
	CBP	-	CREB binding protein
	CCDB	-	Cambridge Crystallographic Data Bank
	cDNA	-	complementary DNA
25	CPU	-	central processing unit
	RAM	-	random access memory
	CRT	-	cathode-ray tube
	DBD	-	DNA binding domain
	DMSO	-	dimethyl sulfoxide
30	DNA	-	deoxyribonucleic acid
	DTT	-	dithiothreitol
	EDTA	-	ethylenediaminetetraacetic acid
	Et <sub>2</sub> O	-	diethyl ether
	FEDs		field emission displays

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	GST	-	glutathione S-transferase
	HEPES	-	N-2-hydroxyethylpiperazine-N'-2-
			ethanesulfonic acid
	kDa	-	kilodalton(s)
5	LBD	-	ligand-binding domain
	LCDs	-	liquid crystal displays
	LED	-	light emitting diode
	MPD	-	methyl-pentanediol
	MCAR	-	mouse constitutive androstane receptor
10	MIR	-	multiple isomorphous replacement
	MPD	-	methyl pentanediol
	N-COR	-	nuclear co-repressor
	NDP	-	nucleotide diphosphate
	NR	-	nuclear receptor
15	nt	-	nucleotide(s)
	NTP	-	nucleotide triphosphate
	PAGE	-	polyacrylamide gel electrophoresis
	PCR	-	polymerase chain reaction
	PEG	-	polyethylene glycol
20	pl	-	isoelectric point
	PXR	-	pregnane X receptor
	PBREM	-	phenobarbital-responsive enhancer module
	RAR	-	retinoic acid receptor
	RAREs	-	retinoic acid response elements
25	rCAR	-	rat constitutive androstane receptor
	RUBISCO	-	ribulose bisphosphate carboxylase
	RXR	-	retinoid X receptor
	SDS	-	sodium dodecyl sulfate
	SDS-PAGE	-	sodium dodecyl sulfate polyacrylamide gel
30			electrophoresis
	SMRT	-	silencing mediator for retinoid and thyroid
			receptors

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SRC-1 - steroid receptor coactivator-1
SR - steroid receptor
TFA - trifluoroacetic acid
TMV - tobacco mosaic virus

TR - thyroid receptor

VDR - vitamin D receptor

Amino Acid Abbreviations, Codes, and Functionally Equivalent Codons

	Amino Acid	3-Letter	1-Letter	Codons
10	Alanine	Ala	Α	GCA GCC GCG GCU
	Arginine	Arg	R	AGA AGG CGA CGC CGG CGU
	Asparagine	Asn	N	AAC AAU
	Aspartic Acid	Asp	D	GAC GAU
	Cysteine	Cys	С	UGC UGU
15	Glutamic acid	Glu	E	GAA GAG
	Glutamine	Gln	Q	CAA CAG
	Glycine	Gly	G	GGA GGC GGG GGU
	Histidine	His	Н	CAC CAU
	Isoleucine	lle	1	AUA AUC AUU
20	Leucine	Leu	L	UUA UUG CUA CUC CUG CUU
	Lysine	Lys	K	AAA AAG
	Methionine	Met	M	AUG
	Phenylalanine	Phe	F	UUC UUU
	Proline	Pro	Р	CCA CCC CCG CCU
25	Serine	Ser	S	ACG AGU UCA UCC UCG UCU
	Threonine	Thr	T	ACA ACC ACG ACU
	Tryptophan	Trp	W	UGG
	Tyrosine	Tyr	Y	UAC UAU
	Valine	Val	V	GUA GUC GUG GUU

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#### **Background**

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The constitutive androstane receptor (CAR; Unified Nomenclature Committee designation NR1I3) was isolated in 1994 by screening a human liver library with a degenerate oligonucleotide probe based on the P box region (Baes et al., 1994). CAR was subsequently shown to be a heterodimer partner for RXR that acts as a specific, retinoid-independent activator of a subset of retinoic acid response elements (RAREs). The mouse CAR homologue was also isolated in 1994 (Honkakoski et al., 1998). Mouse CAR studies showed that RXR and CAR bind to a site in the phenobarbital-responsive enhancer module (PBREM) of the cytochrome P-450 Cyp2b10 gene in response to phenobarbital induction. Expression of RXR and CAR in mammalian cell lines activated PBREM, indicating that a CAR-RXR heterodimer is a trans-acting factor for the mouse Cyp2b10 gene. These studies were the first to indicate that CAR might play a role in response to xenobiotics.

The ability to respond to a wide range of potentially toxic chemicals is essential in a complex environment. Evidence is accumulating that CAR and its closest mammalian homologue, the pregnane X receptor (PXR; Unified Nomenclature Committee designation NR1I2), evolved to detect xenobiotics as part of the body's detoxification machinery (Waxman, 1999). Both receptors are highly expressed in the liver and intestine and both regulate the expression of specific detoxification genes. PXR and CAR regulate genes whose protein products are involved in the hydroxylation (phase I), conjugation (phase II), and transport of xenobiotics (phase III). CAR is activated by some of the same ligands as PXR (Moore et al., 2000), regulates at least partially overlapping sets of genes (e.g. CYP3A and CYP2B; Xie et al., 2000a), and can signal through the same response elements (Goodwin et al., 2001; Handschin et al., 2001).

Despite these similarities, CAR differs from PXR in several respects. CAR ligand binding has been shown to be more restricted than that of PXR (Moore *et al.*, 2000). Furthermore, CAR displays a high basal level of activity relative to PXR that can be reduced by the binding of either naturally

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occurring androstanes or xenobiotics such as clotrimazole (Baes et al., 1994; Moore et al., 2000). Finally, CAR displays fundamental differences from PXR with regard to its cellular regulation. In mouse primary hepatocytes and in mouse liver in vivo, CAR is cytoplasmic in the naïve state and translocates to the nucleus upon activation (Kawamoto et al., 1999), a process thought to be regulated in part by dephosphorylation of the receptor (Honkakoski et al., 1998). Induction of CAR nuclear translocation does not necessarily depend upon ligand-binding, as phenobarbital has been shown to be an activator of CAR in vivo and in hepatocytes, but does not appear to interact directly with the CAR ligand-binding domain (Moore et al., 2000). Thus, CAR has a high basal level of transcriptional activity even in the absence of an exogenous ligand. An important goal of future efforts will be to further differentiate the physical and functional properties of CAR from PXR, and to ultimately distinguish the unique physiological role of CAR.

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Towards this goal, the CAR gene has recently been "knocked-out" by targeted gene disruption (Xie et al., 2000b). The loss of CAR expression did not result in any overt phenotype. Homozygous CAR-1- animals were born at the expected Mendelian frequency, and both male and female CAR-deficient animals were fertile. It was further demonstrated that the nuclear receptor CAR mediates the Cyp2b10 gene response evoked by phenobarbital-like inducers, as well as by the more potent TCPOBOP compound (Xie et al., 2000b). When challenged, these animals showed decreased metabolism of the classic CYP substrate zoxazolamine and a complete loss of the liver hypertrophic and hyperplastic responses to these compounds. These experiments were thus consistent with the notion that at least one aspect of the physiological role of CAR involves xenobiotic metabolism.

Further insight into CAR is expected to be gleaned from CAR structural studies. The availability of the CAR structure will allow an understanding of ligand modulation of CAR activity and will facilitate the design of novel CAR ligands. The present invention addresses these and other needs in the art.

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#### Summary of the Invention

The present invention provides a crystalline form comprising a substantially pure constitutive androstane receptor (CAR) ligand-binding domain polypeptide. In one embodiment, the crystalline form comprises a substantially pure constitutive androstane receptor (CAR) ligand-binding domain polypeptide in complex with a ligand. In one embodiment, a ligand is 2-(benzhydrylamino)-1-(2-phenylethyl)-1H-benzimidazole-6-carboxamide.

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The present invention also provides a method of generating a crystalline form comprising a constitutive androstane receptor (CAR) ligand-binding domain polypeptide in complex with a ligand, the method comprising:

(a) incubating a solution comprising a constitutive androstane receptor (CAR) ligand-binding domain and a ligand with an equal volume of reservoir; and (b) crystallizing the constitutive androstane receptor (CAR) ligand-binding domain polypeptide and ligand using the hanging drop method, whereby a crystalline form of a constitutive androstane receptor (CAR) ligand-binding domain polypeptide in complex with a ligand is generated. Also provided is a crystalline form formed by the above-recited method. In one embodiment, a ligand is 2-(benzhydrylamino)-1-(2-phenylethyl)-1H-benzimidazole-6-carboxamide.

The present invention also provides a method of designing a chemical compound that modulates the biological activity of a target constitutive androstane receptor (CAR) polypeptide. In one embodiment, the method comprises: obtaining one or more three-dimensional structures for the ligand-binding domain (LBD) of constitutive androstane receptor (CAR) in a repressed conformation, and one or more three-dimensional structures of the LBD of constitutive androstane receptor (CAR) in an activated conformation; rotating and translating the three-dimensional structures as rigid bodies so as to superimpose corresponding backbone atoms of a core region of the constitutive androstane receptor (CAR) LBD; comparing one or both of: (i) the superimposed three-dimensional structures to identify volume near the ligand-binding pocket of the constitutive androstane receptor (CAR) LBD that is available to a ligand in the one or more activated structures, or in one or more

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repressed structures, but that is not available to the ligand in one or more structures of the opposite class; and (ii) the superimposed three-dimensional structures to identify interactions that a ligand could make in one or more of the activated structures, or in one or more of the repressed structures, but which the ligand could not make in one or more structures of the opposite class; and designing a chemical compound that occupies the volume, makes the interaction, or both occupies the volume and makes the interaction.

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Optionally the method further comprises synthesizing the designed chemical compound; and testing the designed chemical compound in a biological assay to determine whether it acts as a ligand of constitutive androstane receptor (CAR) with an effect on constitutive androstane receptor (CAR) biological activities, whereby a ligand of a constitutive androstane receptor (CAR) polypeptide is designed.

In another embodiment, the volume or interaction is available in one or more of the repressed structures of constitutive androstane receptor (CAR). but not available in one or more of the activated structures of constitutive androstane receptor (CAR). In another embodiment, the method further comprises designing a chemical compound that promotes the binding of corepressor to the constitutive androstane receptor (CAR) LBD by making direct favorable interactions with the co-repressor. In another embodiment, the method further comprises designing a chemical compound that reduces binding of a co-repressor to the constitutive androstane receptor (CAR) LBD by making direct unfavorable interactions with the co-repressor. In another embodiment, the method further comprises designing a chemical compound that promotes coactivator binding by displacing an AF2 helix of the constitutive androstane receptor (CAR) LBD and making direct favorable interactions with a coactivator, where the designing allows for an expected movement of the coactivator within a coactivator/co-repressor binding pocket. In yet another embodiment, the method further comprises designing a chemical compound by considering a known agonist of the constitutive androstane receptor (CAR) and adding a substituent that protrudes into the volume identified in step (c) or that makes a desired interaction.

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The present invention also provides a binding site in a human constitutive androstane receptor (CAR) polypeptide for a constitutive androstane receptor ligand, wherein the ligand is in van der Waals, hydrogen binding, or van der Waals and hydrogen binding contact with at least one residue of the human constitutive androstane receptor polypeptide.

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The present invention also provides a complex of a human constitutive androstane receptor (CAR) ligand-binding domain and a ligand, wherein the ligand is in van der Waals, hydrogen bonding, or both van der Waals and hydrogen bonding contact with at least one of the following residues of the human constitutive androstane receptor polypeptide: Phe161, Ile164, Asn165, Val199, His203, Phe217, Trp224, Thr225, Ile226, Asp228, Gly229, Gln234, Phe238, Leu239, Leu242, Phe243, Tyr326, Met339, Met340.

The present invention also provides a crystal of a complex of a human constitutive androstane receptor (CAR) ligand-binding domain and a ligand, wherein the ligand is in van der Waals, hydrogen bonding, or both van der Waals and hydrogen bonding contact with at least one of the following residues of the human constitutive androstane receptor polypeptide: Phe161, lle164, Asn165, Val199, His203, Phe217, Trp224, Thr225, lle226, Asp228, Gly229, Gln234, Phe238, Leu239, Leu242, Phe243, Tyr326, Met339, Met340. In one embodiment, the constitutive androstane receptor is a human constitutive androstane receptor and the crystal has the following physical measurements: space group  $P2_12_12_1$ , and unit cell: a = 83.0 angstroms, b = 116.8 angstroms, c = 131.9 angstroms, and a = 8 = y = 90 degrees.

The present invention also provides a method for designing a ligand of a constitutive androstane receptor (CAR) polypeptide, the method comprising: (a) forming a complex of a compound bound to the constitutive androstane receptor (CAR) polypeptide; (b) determining a structural feature of the complex formed in (a); wherein the structural feature is of a binding site for the compound; and (c) using the structural feature determined in (b) to design a ligand of a constitutive androstane receptor (CAR) polypeptide capable of binding to the binding site of the present invention. In one embodiment, the

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method of the present invention further comprises using a computer-based model of the complex formed in (a) in designing the ligand.

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The present invention also provides a method of designing a ligand that selectively modulates the activity of a constitutive androstane receptor (CAR) polypeptide, the method comprising: (a) evaluating a three-dimensional structure of a crystallized constitutive androstane receptor (CAR) ligandbinding domain polypeptide in complex with a ligand; and (b) synthesizing a potential ligand based on the three-dimensional structure of the crystallized constitutive androstane receptor (CAR) catalytic polypeptide in complex with a ligand, whereby a ligand that selectively modulates the activity of a constitutive androstane receptor (CAR) polypeptide is designed. embodiment, the constitutive androstane receptor (CAR) ligand-binding domain polypeptide comprises the amino acid sequence of SEQ ID NO: 4. In one embodiment, the crystalline form is such that the three-dimensional structure of the crystallized constitutive androstane receptor (CAR) ligandbinding domain polypeptide in complex with a ligand can be determined to a resolution of about 2.15 Å or better. In one embodiment, the method further comprises contacting a constitutive androstane receptor (CAR) ligand-binding domain polypeptide with the potential ligand and a ligand; and assaying the constitutive androstane receptor (CAR) ligand-binding domain polypeptide for binding of the potential ligand, for a change in activity of the constitutive androstane receptor (CAR) ligand-binding domain polypeptide, or both. In one embodiment, the ligand is 2-(benzhydrylamino)-1-(2-phenylethyl)-1Hbenzimidazole-6-carboxamide.

The present invention also provides a method of screening a plurality of compounds for a ligand of a constitutive androstane receptor (CAR) ligand-binding domain polypeptide, the method comprising: (a) providing a library of test samples; (b) contacting a crystalline form comprising a constitutive androstane receptor (CAR) polypeptide in complex with a ligand with each test sample; (c) detecting an interaction between a test sample and the crystalline constitutive androstane receptor (CAR) polypeptide in complex with a ligand; (d) identifying a test sample that interacts with the crystalline

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constitutive androstane receptor (CAR) polypeptide in complex with a ligand; and (e) isolating a test sample that interacts with the crystalline constitutive androstane receptor (CAR) polypeptide in complex with a ligand, whereby a plurality of compounds is screened for a ligand of a constitutive androstane receptor (CAR) ligand-binding domain polypeptide. In one embodiment, the CAR polypeptide comprises a CAR ligand-binding domain. In another embodiment, the CAR polypeptide is a human CAR polypeptide. In yet another embodiment, the CAR polypeptide comprises the amino acid sequence of SEQ ID NO: 4. In one embodiment, the library of test samples is bound to a substrate. In another embodiment, the library of test samples is synthesized directly on a substrate. In one embodiment, the ligand is 2-(benzhydrylamino)-1-(2-phenylethyl)-1H-benzimidazole-6-carboxamide,

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The present invention also provides a method for identifying a constitutive androstane receptor (CAR) ligand, the method comprising: (a) providing atomic coordinates of a constitutive androstane receptor (CAR) ligand-binding domain in complex with a ligand to a computerized modeling system; and (b) modeling a ligand that fits spatially into the binding pocket of the constitutive androstane receptor (CAR) ligand-binding domain to thereby identify a constitutive androstane receptor (CAR) ligand. In one embodiment, the method further comprises identifying in an assay for constitutive androstane receptor (CAR)-mediated activity a modeled ligand that increases or decreases the activity of the constitutive androstane receptor (CAR). In one embodiment, the CAR is a human CAR. In one embodiment, the CAR ligand-binding domain comprises the amino acid sequence of SEQ ID NO: 4. In one embodiment, the ligand is 2-(benzhydrylamino)-1-(2-phenylethyl)-1H-benzimidazole-6-carboxamide.

The present invention also provides a method of identifying a constitutive androstane receptor (CAR) ligand that selectively binds a constitutive androstane receptor (CAR) polypeptide compared to other polypeptides, the method comprising: (a) providing atomic coordinates of a constitutive androstane receptor (CAR) ligand-binding domain in complex with a ligand to a computerized modeling system; and (b) modeling a ligand that

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fits into the binding pocket of a constitutive androstane receptor (CAR) ligand-binding domain and that interacts with residues of a constitutive androstane receptor (CAR) ligand-binding domain that are conserved among constitutive androstane receptor (CAR) subtypes to thereby identify a constitutive androstane receptor (CAR) ligand that selectively binds a constitutive androstane receptor (CAR) polypeptide compared to other polypeptides. In one embodiment, the method further comprises identifying in a biological assay for constitutive androstane receptor (CAR) activity a modeled ligand that selectively binds to said constitutive androstane receptor (CAR) and increases or decreases the activity of the constitutive androstane receptor (CAR). In one embodiment, the CAR ligand-binding domain comprises the amino acid sequence shown in SEQ ID NO: 4. In one embodiment, the ligand is 2-(benzhydrylamino)-1-(2-phenylethyl)-1H-benzimidazole-6-carboxamide.

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The present invention also provides a method of designing a ligand of a constitutive androstane receptor (CAR) polypeptide, the method comprising: (a) selecting a candidate constitutive androstane receptor (CAR) ligand; (b) determining which amino acid or amino acids of a constitutive androstane receptor (CAR) polypeptide interact with the ligand using a three-dimensional model of a crystallized protein, the model comprising a constitutive androstane receptor (CAR) ligand-binding domain in complex with a ligand; (c) identifying in a biological assay for constitutive androstane receptor (CAR) activity a degree to which the ligand modulates the activity of the constitutive androstane receptor (CAR) polypeptide; (d) selecting a chemical modification of the ligand wherein the interaction between the amino acids of the constitutive androstane receptor (CAR) polypeptide and the ligand is predicted to be modulated by the chemical modification; (e) synthesizing a ligand having the chemical modified to form a modified ligand; (f) contacting the modified ligand with the constitutive androstane receptor (CAR) polypeptide; (g) identifying in a biological assay for constitutive androstane receptor (CAR) activity a degree to which the modified ligand modulates the biological activity of the constitutive androstane receptor (CAR) polypeptide; and (h) comparing the biological activity of the constitutive androstane

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receptor (CAR) polypeptide in the presence of modified ligand with the biological activity of the constitutive androstane receptor (CAR) polypeptide in the presence of the unmodified ligand, whereby a ligand of a constitutive androstane receptor (CAR) polypeptide is designed. In one embodiment, wherein the method further comprises repeating steps (a) through (f), if the biological activity of the constitutive androstane receptor (CAR) polypeptide in the presence of the modified ligand varies from the biological activity of the constitutive androstane receptor (CAR) polypeptide in the presence of the unmodified ligand.

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The present invention also provides a crystallized, recombinant polypeptide comprising: (a) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (b) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (c) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of constitutive androstane receptor (CAR); wherein the polypeptide of (a), (b) or (c) is in crystal form. In one embodiment, the crystallized, recombinant polypeptide diffracts X-rays to a resolution of about 2.5 Å or better. In another embodiment, the polypeptide is labeled with seleno-methionine.

The present invention also provides a method for designing a modulator for the prevention or treatment of a disease or disorder, comprising:

(a) providing a three-dimensional structure for a crystallized, recombinant polypeptide; (b) identifying a potential modulator for the prevention or treatment of a disease or disorder by reference to the three-dimensional structure; (c) contacting a polypeptide or a constitutive androstane receptor (CAR) with the potential modulator; and (d) assaying the activity of the polypeptide after contact with the modulator, wherein a change in the activity of the polypeptide indicates that the modulator can be useful for prevention or treatment of a disease or disorder.

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The present invention also provides a method for obtaining structural information of a crystallized polypeptide, the method comprising: (a) crystallizing a recombinant polypeptide, wherein the polypeptide comprises: (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); and wherein the crystallized polypeptide is capable of diffracting X-rays to a resolution of 2.5 Å or better; and (b) analyzing the crystallized polypeptide by X-ray diffraction to determine the three-dimensional structure of at least a portion of the crystallized polypeptide. In one embodiment, the three-dimensional structure of the portion of the crystallized polypeptide is determined to a resolution of 2.5 Å or better.

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The present invention also provides a method for identifying a druggable region of a polypeptide, the method comprising: (a) obtaining crystals of a polypeptide comprising (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR), such that the three dimensional structure of the crystallized polypeptide can be determined to a resolution of 2.5 Å or better; (b) determining the three dimensional structure of the crystallized polypeptide using X-ray diffraction; and (c) identifying a druggable region of the crystallized polypeptide based on the three-dimensional structure of the crystallized polypeptide. embodiment, the druggable region is an active site. In another embodiment, the druggable region is on the surface of the polypeptide.

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The present invention also provides a crystalline human constitutive androstane receptor (CAR) comprising a crystal having unit cell dimensions a = 83.0 Å; b = 116.8 Å; c = 131.9 Å;  $\alpha = \beta = \gamma = 90^{\circ}$ ; with an orthorhombic space group P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub> and 4 molecules per asymmetric unit.

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The present invention also provides a crystallized polypeptide comprising: (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); wherein the crystal has a P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub> space group.

The present invention also provides a crystallized polypeptide comprising a structure of a polypeptide that is defined by a substantial portion of the atomic coordinates set forth in Table 2 or Table 3.

The present invention also provides a method for determining the crystal structure of a homolog of a polypeptide, the method comprising: (a) providing the three dimensional structure of a first crystallized polypeptide comprising (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); (b) obtaining crystals of a second polypeptide comprising an amino acid sequence that is at least 70% identical to the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4, such that the three dimensional structure of the second crystallized polypeptide can be determined to a resolution of 2.5 Å or better; and (c) determining the three dimensional structure of the second crystallized polypeptide by X-ray crystallography based on the atomic coordinates of the

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three dimensional structure provided in step (a). In one embodiment, the atomic coordinates for the second crystallized polypeptide have a root mean square deviation from the backbone atoms of the first polypeptide of not more than 1.5 Å for all backbone atoms shared in common with the first polypeptide and the second polypeptide.

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The present invention also provides a method for homology modeling a homolog of human constitutive androstane receptor (CAR), comprising: (a) aligning the amino acid sequence of a homolog of human constitutive androstane receptor (CAR) with an amino acid sequence of SEQ ID NO: 2 or SEQ ID NO: 4 and incorporating the sequence of the homolog of human CAR into a model of human constitutive androstane receptor (CAR) derived from structure coordinates as listed in Table 2 or Table 3 to yield a preliminary model of the homolog of human CAR; (b) subjecting the preliminary model to energy minimization to yield an energy minimized model; (c) remodeling regions of the energy minimized model where stereochemistry restraints are violated to yield a final model of the homolog of human constitutive androstane receptor (CAR).

The present invention also provides a method for obtaining structural information about a molecule or a molecular complex of unknown structure comprising: (a) crystallizing the molecule or molecular complex; (b) generating an X-ray diffraction pattern from the crystallized molecule or molecular complex; (c) applying at least a portion of the structure coordinates set forth in Table 2 or Table 3 to the X-ray diffraction pattern to generate a three-dimensional electron density map of at least a portion of the molecule or molecular complex whose structure is unknown.

The present invention also provides a method for attempting to make a crystallized complex comprising a polypeptide and a modulator having a molecular weight of less than 5 kDa, the method comprising: (a) crystallizing a polypeptide comprising (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that

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hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); such that crystals of the crystallized polypeptide will diffract X-rays to a resolution of 5 Å or better; and (b) soaking the crystals in a solution comprising a potential modulator having a molecular weight of less than 5 kDa.

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The present invention also provides a method for incorporating a potential modulator in a crystal of a polypeptide, comprising placing a hexagonal crystal of human constitutive androstane receptor (CAR) having unit cell dimensions a = 83.0 Å; b = 116.8 Å; c = 131.9 Å,  $a = b = g = 90^{\circ}$ , with an orthorhombic space group P212121, in a solution comprising the potential modulator.

The present invention also provides a computer readable storage medium comprising digitally encoded structural data, wherein the data comprises structural coordinates as listed in Table 2 or Table 3 for the backbone atoms of at least about six amino acid residues from a druggable region of human constitutive androstane receptor (CAR).

The present invention also provides a scalable three-dimensional configuration of points, at least a portion of the points derived from some or all of the structure coordinates as listed in Table 2 or Table 3 for a plurality of amino acid residues from a druggable region of human constitutive androstane receptor (CAR). In one embodiment, the structure coordinates as listed in Table 2 or Table 3 for the backbone atoms of at least about five amino acid residues from a druggable region of human constitutive androstane receptor (CAR) are used to derive part or all of the portion of points. In another embodiment, the structure coordinates as listed in Table 2 or Table 3 for the backbone and optionally the side chain atoms of at least about ten amino acid residues from a druggable region of human constitutive androstane receptor (CAR) are used to derive part or all of the portion of points. In another embodiment, the structure coordinates as listed in Table 2 or Table 3 for the backbone atoms of at least about fifteen amino acid residues from a druggable region of human constitutive androstane receptor

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(CAR) are used to derive part or all of the portion of points. In another embodiment, substantially all of the points are derived from structure coordinates as listed in Table 2 or Table 3. In still another embodiment, the structure coordinates as listed in Table 2 or Table 3 for the atoms of the amino acid residues from any of the above-described druggable regions of human constitutive androstane receptor (CAR) are used to derive part or all of the portion of points.

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The present invention also provides a scalable three-dimensional configuration of points, comprising points having a root mean square deviation of less than about 1.5 Å from the three dimensional coordinates as listed in Table 2 or Table 3 for the backbone atoms of at least five amino acid residues, wherein the five amino acid residues are from a druggable region of human constitutive androstane receptor (CAR). In one embodiment, any point-to-point distance, calculated from the three dimensional coordinates as listed in Table 2 or Table 3, between one of the backbone atoms for one of the five amino acid residues and another backbone atom of a different one of the five amino acid residues is not more than about 10 Å.

The present invention also provides a scalable three-dimensional configuration of points comprising points having a root mean square deviation of less than about 1.5 Å from the three dimensional coordinates as listed in Table 2 or Table 3 for the atoms of the amino acid residues from any of the above-described druggable regions of human constitutive androstane receptor (CAR).

The present invention also provides a computer readable storage medium comprising digitally encoded structural data, wherein the data comprise the identity and three-dimensional coordinates as listed in Table 2 or Table 3 for the atoms of the amino acid residues from any of the above-described druggable regions of human constitutive androstane receptor (CAR).

The present invention also provides a scalable three-dimensional configuration of points, wherein the points have a root mean square deviation of less than about 1.5 Å from the three dimensional coordinates as listed in

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Table 2 or Table 3 for the atoms of the amino acid residues from any of the above-described druggable regions of human constitutive androstane receptor (CAR), wherein up to one amino acid residue in each of the regions can have a conservative substitution thereof.

The present invention also provides a scalable three-dimensional configuration of points derived from a druggable region of a polypeptide, wherein the points have a root mean square deviation of less than about 1.5 Å from the three dimensional coordinates as listed in Table 2 or Table 3 for the backbone atoms of at least ten amino acid residues that participate in the intersubunit contacts of human constitutive androstane receptor (CAR).

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The present invention also provides a computer-assisted method for identifying an inhibitor of the activity of human constitutive androstane receptor (CAR), comprising: (a) supplying a computer modeling application with a set of structure coordinates as listed in Table 2 or Table 3 for the atoms of the amino acid residues from any of the above-described druggable regions of human constitutive androstane receptor (CAR) so as to define part or all of a molecule or complex; (b) supplying the computer modeling application with a set of structure coordinates of a chemical entity; and (c) determining whether the chemical entity is expected to bind to or interfere with the molecule or complex. In one embodiment, determining whether the chemical entity is expected to bind to or interfere with the molecule or complex comprises performing a fitting operation between the chemical entity and a druggable region of the molecule or complex, followed by computationally analyzing the results of the fitting operation to quantify the association between the chemical entity and the druggable region. In one embodiment, the method further comprises screening a library of chemical entities.

The present invention also provides a computer-assisted method for designing an inhibitor of constitutive androstane receptor (CAR) activity comprising: (a) supplying a computer modeling application with a set of structure coordinates having a root mean square deviation of less than about 1.5 Å from the structure coordinates as listed in Table 2 or Table 3 for the atoms of the amino acid residues from any of the above-described druggable

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regions of human constitutive androstane receptor (CAR) so as to define part or all of a molecule or complex; (b) supplying the computer modeling application with a set of structure coordinates for a chemical entity; (c) evaluating the potential binding interactions between the chemical entity and the molecule or complex; (d) structurally modifying the chemical entity to yield a set of structure coordinates for a modified chemical entity; and (e) determining whether the modified chemical entity is an inhibitor expected to bind to or interfere with the molecule or complex, wherein binding to or interfering with the molecule or molecular complex is indicative of potential inhibition of constitutive androstane receptor (CAR) activity. embodiment, determining whether the modified chemical entity is an inhibitor expected to bind to or interfere with the molecule or complex comprises performing a fitting operation between the chemical entity and the molecule or complex, followed by computationally analyzing the results of the fitting operation to evaluate the association between the chemical entity and the molecule or complex. In another embodiment, the set of structure coordinates for the chemical entity is obtained from a chemical library.

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The present invention also provides a computer-assisted method for designing an inhibitor of constitutive androstane receptor (CAR) activity de novo comprising: (a) supplying a computer modeling application with a set of three-dimensional coordinates derived from the structure coordinates as listed in Table 2 or Table 3 for the atoms of the amino acid residues from any of the above-described druggable regions of human constitutive androstane receptor (CAR) so as to define part or all of a molecule or complex; (b) computationally building a chemical entity represented by a set of structure coordinates; and (c) determining whether the chemical entity is an inhibitor expected to bind to or interfere with the molecule or complex, wherein binding to or interfering with the molecule or complex is indicative of potential inhibition of constitutive androstane receptor (CAR) activity. In one embodiment, determining whether the chemical entity is an inhibitor expected to bind to or interfere with the molecule or complex comprises performing a fitting operation between the chemical entity and a druggable region of the

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molecule or complex, followed by computationally analyzing the results of the fitting operation to quantify the association between the chemical entity and the druggable region.

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The present invention also provides a method for identifying a potential modulator for the prevention or treatment of a disease or disorder, the method comprising: (a) providing the three dimensional structure of a crystallized polypeptide comprising: (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); (b) obtaining a potential modulator for the prevention or treatment of a disease or disorder based on the three dimensional structure of the crystallized polypeptide; (c) contacting the potential modulator with a second polypeptide comprising: (i) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (ii) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4: or (iii) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); which second polypeptide can optionally be the same as the crystallized polypeptide; and (d) assaying the activity of the second polypeptide, wherein a change in the activity of the second polypeptide indicates that the compound can be useful for prevention or treatment of a disease or disorder.

The present invention also provides a method for designing a candidate modulator for screening for inhibitors of a polypeptide, the method comprising: (a) providing the three dimensional structure of a druggable region of a polypeptide comprising (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least

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about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); and (b) designing a candidate modulator based on the three dimensional structure of the druggable region of the polypeptide.

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The present invention also provides a method for identifying a potential modulator of a polypeptide from a database, the method comprising: (a) providing the three-dimensional coordinates for a plurality of the amino acids of a polypeptide comprising (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); (b) identifying a druggable region of the polypeptide; and (c) selecting from a database at least one potential modulator comprising three dimensional coordinates which indicate that the modulator can bind or interfere with the druggable region. In one embodiment, the modulator is a small molecule.

The present invention also provides a method for preparing a potential modulator of a druggable region contained in a polypeptide, the method comprising: (a) using the atomic coordinates for the backbone atoms of at least about six amino acid residues from a polypeptide of SEQ ID NO: 4, with a root mean square deviation from the backbone atoms of the amino acid residues of not more than 1.5 Å, to generate one or more three-dimensional structures of a molecule comprising a druggable region from the polypeptide; (b) employing one or more of the three dimensional structures of the molecule to design or select a potential modulator of the druggable region; and (c) synthesizing or obtaining the modulator.

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The present invention also provides an apparatus for determining whether a compound is a potential modulator of a polypeptide, the apparatus comprising: (a) a memory that comprises: (i) the three dimensional coordinates and identities of at least about fifteen atoms from a druggable region of a polypeptide comprising (1) an amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; (2) an amino acid sequence having at least about 95% identity with the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; or (3) an amino acid sequence encoded by a polynucleotide that hybridizes under stringent conditions to the complementary strand of a polynucleotide having SEQ ID NO: 1 or SEQ ID NO: 3 and has at least one biological activity of human constitutive androstane receptor (CAR); (ii) executable instructions; and (b) a processor that is capable of executing instructions to: (i) receive three-dimensional structural information for a candidate modulator; (ii) determine if the three-dimensional structure of the candidate modulator is complementary to the three dimensional coordinates of the atoms from the druggable region; and (iii) output the results of the determination.

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The present invention also provides a method for making an inhibitor of constitutive androstane receptor (CAR) activity, the method comprising chemically or enzymatically synthesizing a chemical entity to yield an inhibitor of constitutive androstane receptor (CAR) activity, the chemical entity having been identified during a computer-assisted process comprising supplying a computer modeling application with a set of structure coordinates of a molecule or complex, the molecule or complex comprising at least a portion of at least one druggable region from human constitutive androstane receptor (CAR); supplying the computer modeling application with a set of structure coordinates of a chemical entity; and determining whether the chemical entity is expected to bind or to interfere with the molecule or complex at a druggable region, wherein binding to or interfering with the molecule or complex is indicative of potential inhibition of constitutive androstane receptor (CAR) activity.

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The present invention also provides a computer readable storage medium comprising digitally encoded data, wherein the data comprises structural coordinates for a druggable region that is structurally homologous to the structure coordinates as listed in Table 2 or Table 3 for a druggable region of human constitutive androstane receptor (CAR).

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The present invention also provides a computer readable storage medium comprising digitally encoded structural data, wherein the data comprise a majority of the three-dimensional structure coordinates as listed in Table 2 or Table 3. In one embodiment, the computer readable storage medium further comprises the identity of the atoms for the majority of the three-dimensional structure coordinates as listed in Table 2 or Table 3. In another embodiment, the data comprise substantially all of the three-dimensional structure coordinates as listed in Table 2 or Table 3.

The present invention also provides a method for building a model for an activated conformation of a constitutive androstane receptor (CAR), the method comprising: (a) employing coordinates for CAR residues 107 to 332 as shown in Table 2; (b) rotating and translating an X-ray structure of the Vitamin D receptor (VDR), so as to superimpose its core backbone atoms onto corresponding atoms from CAR; (c) combining a superimposed VDR AF2 helix, residues 416-423, with residues 107-332 from CAR from step (a), to provide a starting model for residues 107-332 and 341-348 of CAR in the activated conformation; (d) computationally mutating Val418, Leu419, Val421. Phe422 and Gly423 in the VDR AF2 helix to corresponding amino acids in a CAR AF2 helix, wherein the corresponding amino acids in the CAR AF2 helix are Leu343, Gln344, Ile346, Cys347 and Ser348, respectively; and (e) adjusting the conformations of the mutated amino acid side chains in residues 343, 344, and 346-348 of the AF2 helix of CAR to avoid overlaps, wherein the adjusting is accomplished by one of manual manipulation and conformational search and energy minimization. In one embodiment, the method further comprises modeling a CAR AF2 linker region, residues 333-340, by using a computational loop modeling technique.

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Accordingly, it is an object of the present invention to provide a threedimensional structure of the ligand-binding domain of CAR in complex with a ligand. The object is achieved in whole or in part by the present invention.

An object of the invention having been stated hereinabove, other objects will be evident as the description proceeds, when taken in connection with the accompanying Drawings and Examples as described hereinbelow.

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# Brief Description of the Drawings

Figure 1 is a ribbon diagram depicting the secondary structure of CAR LBD bound with ligand. The ligand is shown as ball and stick. Helices are indicated by  $\bf H$  followed by the  $\alpha$  helix number, and  $\beta$ -strands are indicated by  $\bf b$  followed by the  $\beta$ -strand number. The line at the bottom of the figure indicates the scale, and corresponds to 50 angstroms.  $\bf N$  refers to the N-terminus and  $\bf C$  refers to the C-terminus.

Figure 2 is a structure-based sequence alignment of the human, mouse, and rat CAR polypeptides with the human PXR polypeptide and the human VDR polypeptide. The residues that make up the  $\alpha$  helices are boxed with a light gray line and light gray background. The residues that make up the  $\beta$  sheets are boxed with a darker gray line and darker gray background. The residues within 5Å of the ligand are individually boxed with a thin black square box. Conserved residues are indicated in bold type.

Figure 3 depicts the CAR ligand-binding site. CAR amino acids are shown with light and dark gray lines. A ligand is shown in heavy black lines. The hydrogen bonds between CAR amino acids and the ligand are shown with dotted lines. Particular amino acids that are involved in the ligand binding are indicated using one letter code and amino acid number.

Figure 4 is a stick diagram depicting another view of the ligand-binding site. CAR amino acids are shown with light and dark gray lines. A ligand is shown in heavy black lines. The hydrogen bonds between CAR amino acids and the ligand are shown with dotted lines. Particular amino acids that are involved in the ligand binding are indicated using one letter code and amino acid number.

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Figure 5 depicts the CAR binding pocket. Ligand Compound 1 is shown in Van der Walls ball form. The binding pocket is shown as a dotted surface. The protein backbone is shown in ribbon form. The side chains in the binding pocket are shown in ball and stick form.

Figure 6 depicts another view of the ribbon diagram depicting secondary structure of the three-layer sandwich shaped ligand-binding pocket.

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Figure 7 is a schematic diagram of a general strategy for synthesizing ligands that can bind to the CAR LBD. This scheme is described in Example 6, which outlines the synthesis of an exemplary ligand, Compound 1.

# Brief Description of the Sequences in the Sequence Listing

SEQ ID NO: 1 is a DNA sequence encoding a full-length human CAR polypeptide.

SEQ ID NO: 2 is an amino acid sequence of a full-length human CAR polypeptide.

SEQ ID NO: 3 is a DNA sequence encoding human CAR residues 103-340, the ligand-binding domain of CAR polypeptide.

SEQ ID NO: 4 is an amino acid sequence of residues 103-340, the ligand-binding domain of CAR polypeptide.

SEQ ID NO: 5 is a His tag amino acid sequence.

SEQ ID NO: 6 is a DNA sequence of a primer used in combination with the primer of SEQ ID NO: 7 to amplify a DNA fragment encoding amino acid residues 103 - 348 of a human CAR polypeptide. In addition to amplifying these coding nucleotides, the primer also includes sequences that will result in the amplified product (a) encoding a His tag as in SEQ ID NO: 5; and (b) having an Ndel endonuclease restriction site (CATATG) just 5' to the His tagencoding residues.

SEQ ID NO: 7 is a DNA sequence of a primer used in combination with the primer of SEQ ID NO: 6 to amplify a DNA fragment encoding residues 103 - 348 of a human CAR polypeptide. The sequence of this primer includes a BamHI endonuclease restriction site (GGATCC) 3' to the human CAR

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polypeptide coding residues. When this primer is used in combination with the primer of SEQ ID NO: 6, the amplified product will have the following arrangement of features: Ndel site – His tag – nucleotides encoding human CAR amino acids 103 to 348 – BamHI site.

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# **Detailed Description of the Invention**

Until disclosure of the present invention presented herein, the ability to obtain crystalline forms of a CAR LBD, particularly in complex with an antagonist ligand, has not been realized. And until disclosure of the present invention presented herein, a detailed three-dimensional crystal structure of an unliganded CAR polypeptide or a CAR polypeptide in complex with a ligand has not been solved.

In addition to providing structural information, crystalline polypeptides provide other advantages. For example, the crystallization process itself further purifies the polypeptide, and satisfies one of the classical criteria for homogeneity. In fact, crystallization frequently provides unparalleled purification quality, removing impurities that are not removed by other purification methods such as HPLC, dialysis, conventional column chromatography, etc. Moreover, crystalline polypeptides are often stable at ambient temperatures and free of protease contamination and degradation associated with solution storage. Crystalline polypeptides can also be useful as pharmaceutical preparations. Finally, crystallization techniques are generally free of problems such as denaturation associated with other stabilization methods (e.g., lyophilization).

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Once crystallization has been accomplished, crystallographic data provides useful structural information that can assist the design of compounds that can serve as agonists or antagonists, as described herein below. In addition, the crystal structure provides information that can be used to map the molecular surface of the ligand-binding domain of CAR. A small non-peptide molecule designed to mimic portions of this surface could serve as a modulator of CAR activity.

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## <u>I.</u> <u>Definitions</u>

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Before the present proteins, nucleotide sequences, and methods are described, it is understood that this invention is not limited to the particular methodology, protocols, cell lines, vectors, and reagents described, as these can vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to limit the scope of the present invention, the invention being defined by the claims.

Unless defined otherwise, all technical and scientific terms used herein are intended to have their ordinary meanings as understood by one of ordinary skill in the art to which this invention pertains. Although any methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, representative methods, devices, and materials are now described. All publications mentioned herein are incorporated by reference for the purpose of describing the cell lines, vectors, reagents, and methodologies they disclose.

Following long-standing patent law convention, the articles "a" and "an" are used herein to refer to one or to more than one (*i.e.*, to at least one) of the grammatical object of the article. By way of example, "an element" means one element or more than one element.

As used herein, the term "AF2 helix" refers to a short alpha-helix, usually including 5-8 residues, located at the C-terminal end of a LBD sequence, that can usually adopt multiple positions, orientations, and conformations in the structure, and which is involved in binding to coactivators. In the hypothetical activated conformation of CAR, the AF2 helix is expected to include residues 341 to 347. These residues do not adopt an alpha-helical conformation in the structure of CAR bound to Compound 1.

As used herein, the terms "Compound 1" and "Formula (A)" are used interchangeably and refer to 2-(benzhydrylamino)-1-(2-phenylethyl)-1H-benzimidazole-6-carboxamide.

As used herein, the term "AF2 glutamate" refers to a glutamate residue in the AF2 helix that can make hydrogen bond interactions with the exposed

NH groups of the LXXLL-containing peptide from a coactivator if the AF2 helix is in the active position. In CAR, the AF2 glutamate is residue number 345.

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As used herein, the terms "activated", "active conformation", and "activated conformation" of an LBD are used interchangeably and refer to a conformation where the AF2 helix is in the active position, thereby placing the AF2 glutamate residue in a position and orientation that creates a charge clamp that can recruit coactivator peptides. Similarly, the terms "active position of the AF2 helix" and "active conformation of the AF2 helix" are used interchangeably and mean an AF2 helix having a position and/or orientation similar to that of the AF2 helix in the PPARg/SRC-1/rosiglitazone structure of Nolte et al., 1998, allowing the AF2 glutamate residue to make interactions with the exposed NH groups of a coactivator peptide. The position and/or orientation of the AF2 helix in an NR structure can be compared with that of the AF2 helix in another NR structure by rotating and/or translating one structure so as to superimpose the backbone atoms of helices 1 through 10 onto the corresponding atoms of the other structure, where corresponding residues are determined by sequence alignment. If, after superimposition, a majority of the backbone atoms of the core of the AF2 helix lie within 2.0 angstroms of the corresponding atoms from the PAPRg/SRC-1/rosiglitazone structure, then the AF2 helix is defined as being in an active position or active conformation.

Other examples of a nuclear receptor where the AF2 helix is in an "active position" include the X-ray structures of the estrogen receptor  $\alpha$  (ER $\alpha$ ) bound to estradiol (Brzozowski *et al.*, 1997) and diethylstilbesterol (DES) (Shiau *et al.*, 1998). Examples of a nuclear receptor where the AF2 helix is not in an "active position" are the X-ray structures of the estrogen receptor  $\alpha$  (ER $\alpha$ ) bound to raloxifene (Brzozowski *et al.*, 1997) and tamoxifen (Shiau *et al.*, 1998). Binding of a coactivator, and AF2-dependent activation of gene transcription, normally requires that the AF2 helix be in the "active position" (Nolte *et al.*, 1998; Shiau *et al.*, 1998). This creates a "charge-clamp" structure that holds the coactivator in its required position (Nolte *et al.*, 1998).

As used herein, the terms "repressed", "inactive conformation", and "repressed conformation" of an LBD are used interchangeably and refer to a conformation where the AF2 helix is not in the active position, and where the AF2 glutamate residue is not in a position that could create the charge clamp that can recruit coactivator peptides.

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As used herein, the term "agonist" refers to an agent that supplements or potentiates the biological activity of a functional CAR gene or protein, or of a polypeptide encoded by a gene that is up- or down-regulated by a CAR polypeptide and/or a polypeptide encoded by a gene that contains a CAR binding site or response element in its promoter region. An agent is also an agonist when the changes in gene expression, considered over many genes, are similar in direction to those induced by other agents that are commonly regarded as agonists. In one embodiment, an agonist of CAR is an androstane.

As used herein, the term "antagonist" refers to an agent that decreases or inhibits the biological activity of a functional gene or protein (for example, a functional CAR gene or protein), or that supplements or potentiates the biological activity of a naturally occurring or engineered non-functional gene or protein (for example, a non-functional CAR gene or protein). Alternatively, an antagonist can decrease or inhibit the biological activity of a functional gene or polypeptide encoded by a gene that is up- or down-regulated by a CAR polypeptide and/or contains a CAR binding site or response element in its promoter region. An antagonist can also supplement or potentiate the biological activity of a naturally occurring or engineered non-functional gene or polypeptide encoded by a gene that is up- or down-regulated by a CAR polypeptide, and/or contains a CAR binding site or response element in its promoter region. An agent is also an antagonist when the changes in gene expression, considered over many genes, are opposite in direction to those induced by other agents that are commonly regarded as agonists.

As used herein, the terms " $\alpha$ -helix" and "alpha-helix" are used interchangeably and refer to a conformation of a polypeptide chain wherein the polypeptide backbone is wound around the long axis of the molecule in a

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left-handed or right-handed direction, and the R groups of the amino acids protrude outward from the helical backbone, wherein the repeating unit of the structure is a single turn of the helix, which extends about 0.56 nm along the long axis.

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As used herein, the terms "amino acid", "amino acid residue", and "residue" are used interchangeably and refer to an amino acid formed upon chemical digestion (hydrolysis) of a peptide or polypeptide at its peptide linkages. Amino acids can also be synthesized individually or as components of a peptide. In one embodiment, the amino acid residues described herein are in the "L" isomeric form. However, residues in the "D" isomeric form can be substituted for any L-amino acid residue, provided that the desired functional property is retained by the polypeptide. In the context of an amino acid, NH2 refers to the free amino group present at the amino terminus of a polypeptide, although some amino acids can have NH2 groups at other positions in the amino acid. COOH refers to the free carboxy group present at the carboxy terminus of a polypeptide. In keeping with standard polypeptide nomenclature, abbreviations for amino acid residues are presented above. The term "amino acid" is intended to embrace all molecules, whether natural or synthetic, which include both an amino functionality and an acid functionality and capable of being included in a polymer of naturally occurring amino acids. Exemplary amino acids include naturally occurring amino acids; analogs, derivatives and congeners thereof; amino acid analogs having variant side chains; and all stereoisomers of any of the foregoing.

It is noted that amino acid residue sequences represented herein by formulae have a left-to-right orientation in the conventional direction of amino terminus to carboxy terminus. In addition, the terms "amino acid", "amino acid residue", and "residue" are broadly defined to include the amino acids listed in the above table and modified or unusual amino acids. Furthermore, it is noted that a dash at the beginning or end of an amino acid residue sequence indicates a peptide bond to a further sequence of one or more amino acid residues or a covalent bond to an amino-terminal group such as NH<sub>2</sub> or acetyl or to a carboxy-terminal group such as COOH.

As used herein, the terms " $\beta$ -sheet" and "beta-sheet" are used interchangeably and refer to the conformation of a polypeptide chain stretched into an extended zigzag conformation. Portions of polypeptide chains that run "parallel" all run in the same direction. Polypeptide chains that are "antiparallel" run in the opposite direction from the parallel chains or from each other.

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The term "binding" refers to an association, which can be a stable association, between two molecules, *i.e.*, between a polypeptide of the invention and a binding partner, due to, for example, electrostatic, hydrophobic, ionic, and/or hydrogen-bond interactions under physiological conditions.

As used herein, the terms "binding pocket of the CAR ligand-binding domain", "CAR ligand-binding pocket" and "CAR binding pocket" are used interchangeably, and refer to the large cavity within the CAR ligand-binding domain where a ligand (e.g. Compound 1) binds. This cavity can be empty, or can contain water molecules or other molecules from the solvent, or can contain ligand atoms. The "main" binding pocket includes the region of space not occupied by atoms of CAR that is approximately encompassed or bounded by residues Phe132, Phe161, Ile164, Asn165, Thr166, Met168, Val169, Ala198, Val199, Cys202, His203, Leu206, Phe217, Tyr224, Thr225, Ile226, Glu227, Asp228, Gly229, Ala230, Phe234, Phe238, Leu239, Leu242, Phe243, His246, Tyr326, Ile330, Leu336, Ser337, Met339, and Met340. The binding pocket also includes small regions near to and contiguous with the "main" binding pocket that not occupied by atoms of CAR.

As used herein the term "biological activity" refers to any biochemical function of a biological molecule. A biological activity includes, but is not limited to, an interaction with another biological molecule (for example, a polypeptide or a nucleic acid, or a combination thereof). As such, a biological activity results in a biochemical effect including, but not limited to the initiation or inhibition of transcription of a gene.

The term "complex" refers to an association between at least two moieties (i.e. chemical or biochemical) that have an affinity for one another.

Examples of complexes include associations between antigen/antibodies, lectin/avidin, target polynucleotide/probe oligonucleotide, antibody/antiantibody, receptor/ligand, enzyme/ligand, polypeptide/ polypeptide, polypeptide/polynucleotide, polypeptide/co-factor, polypeptide/substrate, polypeptide/inhibitor, polypeptide/small molecule, and the like. "Member of a complex" refers to one moiety of the complex, such as an antigen or ligand. "Protein complex" or "polypeptide complex" refers to a complex comprising at least one polypeptide.

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The term "conserved residue" refers to an amino acid that is a member of a group of amino acids having certain common properties. The term "conservative amino acid substitution" refers to the substitution (conceptually or otherwise) of an amino acid from one such group with a different amino acid from the same group. A functional way to define common properties between individual amino acids is to analyze the normalized frequencies of amino acid changes between corresponding proteins of homologous organisms (Schulz & Schirmer, 1979). According to such analyses, groups of amino acids can be defined where amino acids within a group exchange preferentially with each other, and therefore resemble each other most in their impact on the overall protein structure (Schulz & Schirmer, 1979). Representative examples of sets of amino acid groups defined in this manner include: (i) a charged group, consisting of Glu and Asp, Lys, Arg and His, (ii) a positively-charged group, consisting of Lys, Arg and His, (iii) a negativelycharged group, consisting of Glu and Asp, (iv) an aromatic group, consisting of Phe, Tyr and Trp, (v) a nitrogen ring group, consisting of His and Trp, (vi) a large aliphatic nonpolar group, consisting of Val, Leu and Ile, (vii) a slightlypolar group, consisting of Met and Cys, (viii) a small-residue group, consisting of Ser, Thr, Asp, Asn, Gly, Ala, Glu, Gln and Pro, (ix) an aliphatic group consisting of Val, Leu, IIe, Met and Cys, and (x) a small hydroxyl group consisting of Ser and Thr.

As used herein, the term "DNA segment" refers to a DNA molecule that has been isolated free of total genomic DNA of a particular species. In one embodiment, a DNA segment encoding a CAR polypeptide refers to a nucleic

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acid comprising SEQ ID NO: 1. In another embodiment, a DNA segment encoding a CAR polypeptide refers to a nucleic acid comprising SEQ ID NO: 3. DNA segments can comprise a portion of a recombinant vector, including, for example, a plasmid, a cosmid, a phage, a virus, and the like.

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As used herein, the term "DNA sequence encoding a CAR polypeptide" refers to one or more coding sequences within a particular individual. Moreover, certain differences in nucleotide sequences can exist between individual organisms, which are called alleles. It is possible that such allelic differences might or might not result in differences in amino acid sequence of the encoded polypeptide yet still encode a protein with the same biological activity. As is well known, genes for a particular polypeptide can exist in single or multiple copies within the genome of an individual. Such duplicate genes can be identical or can have certain modifications, including nucleotide substitutions, additions, or deletions, all of which still code for polypeptides having substantially the same activity.

The term "domain", when used in connection with a polypeptide, refers to a specific region within the polypeptide that comprises a particular structure or mediates a particular function. In the typical case, a domain of a polypeptide of the invention is a fragment of the polypeptide. In certain instances, a domain is a structurally stable domain, as evidenced, for example, by mass spectroscopy, or by the fact that a modulator can bind to a druggable region of the domain. In one embodiment, a domain of a CAR polypeptide is a ligand-binding domain. In another embodiment, a domain of a CAR polypeptide is a DNA-binding domain.

The term "druggable region", when used in reference to a polypeptide, nucleic acid, complex and the like, refers to a region of the molecule that is a target or is a likely target for binding a modulator. For a polypeptide, a druggable region generally refers to a region wherein several amino acids of a polypeptide would be capable of interacting with a modulator or other molecule. For a polypeptide or complex thereof, exemplary druggable regions including binding pockets and sites, enzymatic active sites, interfaces between domains of a polypeptide or complex, surface grooves or contours or

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surfaces of a polypeptide or complex which are capable of participating in interactions with another molecule. In certain instances, the interacting molecule is another polypeptide, which can be naturally occurring. In other instances, the druggable region is on the surface of the molecule. In one embodiment, a druggable region of a CAR polypeptide comprises the binding site defined by amino acid residues 103-340. In another embodiment, a druggable region of a CAR polypeptide comprises amino acid residues and surfaces of the CAR polypeptide that interact with a RXR polypeptide during CAR-RXR heterodimer formation. In another embodiment, a druggable region of a CAR polypeptide comprises the AF2 helix. In another embodiment, a druggable region of a CAR polypeptide comprises Glu345. In still another embodiment, a druggable region of a CAR polypeptide comprises a DNA-binding domain.

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Druggable regions can be described and characterized in a number of ways. For example, a druggable region can be characterized by some or all of the amino acids that make up the region, or the backbone atoms thereof, or the side chain atoms thereof (optionally with or without the  $C\alpha$  atoms). Alternatively, in certain instances, the volume of a druggable region corresponds to that of a carbon based molecule of at least about 200 atomic mass units (amu) and often up to about 800 amu. In other instances, it will be appreciated that the volume of such region can correspond to a molecule of at least about 600 amu and often up to about 1600 amu or more.

Alternatively, a druggable region can be characterized by comparison to other regions on the same or other molecules. For example, the term "affinity region" refers to a druggable region on a molecule (such as a polypeptide of the invention) that is present in several other molecules, in so much as the structures of the same affinity regions are sufficiently the same so that they are expected to bind the same or related structural analogs. An example of an affinity region is an ATP-binding site of a protein kinase that is found in several protein kinases (whether or not of the same origin). Another example of an affinity region is a DNA-binding domain: for example, the DNA-binding domain of a CAR polypeptide.

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In contrast to an affinity region, the term "selectivity region" refers to a druggable region of a molecule that can not be found on other molecules, in so much as the structures of different selectivity regions are sufficiently different so that they are not expected to bind the same or related structural analogs. An exemplary selectivity region is a catalytic domain of a protein kinase that exhibits specificity for one substrate. In certain instances, a single modulator can bind to the same affinity region across a number of proteins that have a substantially similar biological function, whereas the same modulator can bind to only one selectivity region of one of those proteins.

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Continuing with examples of different druggable regions, the term "undesired region" refers to a druggable region of a molecule that upon interacting with another molecule results in an undesirable affect. For example, a binding site that oxidizes the interacting molecule and thereby results in increased toxicity for the oxidized molecule can be deemed an "undesired region". Other examples of potential undesired regions include regions that upon interaction with a drug decrease the membrane permeability of the drug, increase the excretion of the drug, or increase the blood brain transport of the drug. It can be the case that, in certain circumstances, an undesired region will no longer be deemed an undesired region because the affect of the region will be favorable, *i.e.*, a drug intended to treat a brain condition would benefit from interacting with a region that resulted in increased blood brain transport, whereas the same region could be deemed undesirable for drugs that were not intended to be delivered to the brain.

When used in reference to a druggable region, the "selectivity" or "specificity' of a molecule such as a modulator to a druggable region can be used to describe the binding between the molecule and a druggable region. For example, the selectivity of a modulator with respect to a druggable region can be expressed by comparison to another modulator, using the respective values of  $K_d$  (*i.e.*, the dissociation constants for each modulator-druggable region complex) or, in cases where a biological effect is observed below the  $K_d$ , the ratio of the respective  $EC_{50}$ 's (*i.e.*, the concentrations that produce

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50% of the maximum response for the modulator interacting with each druggable region).

As used herein, the term "expression" generally refers to the cellular processes by which a biologically active polypeptide is produced. As such, the term "expression" generally includes those cellular processes that begin with transcription and end with the production of a functional polypeptide. As used herein, "expression" is also intended to refer to cellular processes by which a polypeptide is produced that would otherwise be functional except for the presence of mutations in the nucleotide sequence encoding it. Consistent with this usage, "expression" includes, but is not limited to, such processes as transcription, translation, post-translational modification, and transport of a polypeptide.

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A "fusion protein" or "fusion polypeptide" refers to a chimeric protein as that term is known in the art and can be constructed using methods known in In many examples of fusion proteins, there are two different polypeptide sequences, and in certain cases, there can be more. sequences can be linked in frame. A fusion protein can include a domain that is found (albeit in a different protein) in an organism that also expresses the first protein, or it can be an "interspecies", "intergenic", etc. fusion expressed by different kinds of organisms. In various embodiments, the fusion polypeptide can comprise one or more amino acid sequences linked to a first polypeptide. In the case where more than one amino acid sequence is fused to a first polypeptide, the fusion sequences can be multiple copies of the same sequence, or alternatively, can be different amino acid sequences. The fusion polypeptides can be fused to the N-terminus, the C-terminus, or the Nand C-terminus of the first polypeptide. Exemplary fusion proteins include polypeptides comprising a glutathione S-transferase tag (GST-tag), histidine tag (His-tag), an immunoglobulin domain, or an immunoglobulin-binding domain.

As used herein, the term "gene" is used for simplicity to refer to a nucleotide sequence that encodes a protein, a polypeptide, or a peptide. As such, the term "gene" refers to a nucleic acid comprising an open reading

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frame encoding a polypeptide having exon sequences and, optionally, intron sequences. The term "intron" refers to a DNA sequence present in a given gene that is not translated into protein and is generally found between exons. As will be understood by those of skill in the art, this functional term includes both genomic sequences and cDNA sequences. Representative embodiments of such sequences are disclosed herein.

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The term "having substantially similar biological activity", when used in reference to two polypeptides, refers to a biological activity of a first polypeptide which is substantially similar to at least one of the biological activities of a second polypeptide. A substantially similar biological activity means that the polypeptides carry out a similar function, i.e., a similar enzymatic reaction or a similar physiological process, etc. For example, two homologous proteins can have a substantially similar biological activity if they are involved in a similar enzymatic reaction, i.e., they are both kinases which catalyze phosphorylation of a substrate polypeptide, however, they can phosphorylate different regions on the same protein substrate or different substrate proteins altogether. Alternatively, two homologous proteins can also have a substantially similar biological activity if they are both involved in a similar physiological process, i.e., regulation of transcription. For example, two proteins can be transcription factors, however, they can bind to different DNA sequences or bind to different polypeptide interactors. Substantially similar biological activities can also be associated with proteins carrying out a similar structural role, for example, two membrane proteins.

As used herein, the term "interact" refers to detectable interactions between molecules, such as can be detected using, for example, a yeast two-hybrid assay. The term "interact" is also meant to include "binding" interactions between molecules. Interactions include, but are not limited to protein-protein, protein-nucleic acid, and protein-small molecule interactions. These interactions can be in the form of covalent or non-covalent interactions including, but not limited to ionic, hydrogen bonding, and van der Waals interactions.

As used herein, the term "isolated" refers to a nucleic acid substantially free of other nucleic acids, proteins, lipids, carbohydrates, or other materials with which it can be associated, such association being either in cellular material or in a synthesis medium. The term can also be applied to polypeptides, in which case the polypeptide is substantially free of nucleic acids, carbohydrates, lipids, and other undesired polypeptides. The term "isolated polypeptide" refers to a polypeptide, in certain embodiments prepared from recombinant DNA or RNA, or of synthetic origin, or some combination thereof, which (1) is not associated with proteins that it is normally found with in nature, (2) is isolated from the cell in which it normally occurs, (3) is isolated free of other proteins from the same cellular source, (4) is expressed by a cell from a different species, or (5) does not occur in nature.

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The term "isolated nucleic acid" refers to a polynucleotide of genomic, cDNA, or synthetic origin or some combination there of, which (1) is not associated with the cell in which the "isolated nucleic acid" is found in nature, or (2) is operably linked to a polynucleotide to which it is not linked in nature.

The terms "label" or "labeled" refer to incorporation or attachment, optionally covalently or non-covalently, of a detectable marker into a molecule, such as a polypeptide. Various methods of labeling polypeptides are known in the art and can be used. Examples of labels for polypeptides include, but are not limited to the following: radioisotopes, fluorescent labels, heavy atoms, enzymatic labels or reporter genes, chemiluminescent groups, biotinyl groups, predetermined polypeptide epitopes recognized by a secondary reporter (i.e., leucine zipper pair sequences, binding sites for secondary antibodies, metal binding domains, epitope tags). Examples and use of such labels are well known by the skilled artisan. In some embodiments, spacer arms of various lengths can be attached to labels to reduce potential steric hindrance.

The term "mammal" is known in the art, and exemplary mammals include humans, primates, bovines, porcines, canines, felines, and rodents (i.e., mice and rats).

The term "modulation", when used in reference to a functional property or biological activity or process (*i.e.*, enzyme activity or receptor binding), refers to the capacity to up regulate (*i.e.*, activate or stimulate), down regulate (*i.e.*, inhibit or suppress), or otherwise change a quality of such property, activity, or process. In certain instances, such regulation can be contingent on the occurrence of a specific event, such as activation of a signal transduction pathway, and/or can be manifest only in particular cell types.

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The term "modulator" refers to a polypeptide, nucleic acid, macromolecule, complex, molecule, small molecule, compound, species, or the like (naturally-occurring or non-naturally-occurring), or an extract made from biological materials such as bacteria, plants, fungi, or animal cells or tissues, that can be capable of causing modulation. Modulators can be evaluated for potential activity as inhibitors or activators (directly or indirectly) of a functional property, biological activity or process, or combination thereof, (i.e., agonist, partial antagonist, partial agonist, inverse agonist, antagonist, anti-microbial agents, inhibitors of microbial infection or proliferation, and the like) by inclusion in assays. In such assays, many modulators can be screened at one time. The activity of a modulator can be known, unknown, or partially known.

As used herein, the term "molecular replacement" refers to a method that involves generating a preliminary model of the wild-type CAR ligand-binding domain, or a CAR mutant crystal the structure for which coordinates are unknown, by orienting and positioning a molecule the structure for which coordinates are known (e.g., the vitamin D receptor; VDR) within the unit cell of the unknown crystal so as best to account for the observed diffraction pattern of the unknown crystal. Phases can then be calculated from this model and combined with the observed amplitudes to give an approximate Fourier synthesis of the structure the coordinates for which are unknown. This, in turn, can be subjected to any of the several forms of refinement known in the art to provide a final, accurate structure of the unknown crystal (see e.g. Lattman, 1985; Rossmann, 1972). Using the structure coordinates of the ligand-binding domain of CAR provided by this invention, molecular

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replacement can be used to determine the structure coordinates of a crystal of a mutant or of a homologue of the CAR ligand-binding domain, or of a different crystal form of the CAR ligand-binding domain.

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The term "motif" refers to an amino acid sequence that is commonly found in a protein of a particular structure or function. Typically, a consensus sequence is defined to represent a particular motif. The consensus sequence need not be strictly defined and can contain positions of variability, degeneracy, variability of length, etc. The consensus sequence can be used to search a database to identify other proteins that can have a similar structure or function due to the presence of the motif in its amino acid For example, on-line databases can be searched with a sequence. consensus sequence in order to identify other proteins containing a particular motif. Various search algorithms and/or programs can be used, including FASTA, BLAST, or ENTREZ. FASTA and BLAST are available as a part of the GCG sequence analysis package (Accelrys, Inc., San Diego, California, United States of America). ENTREZ is available through the National Center for Biotechnology Information, National Library of Medicine, National Institutes of Health, Bethesda, Maryland, United States of America.

As used herein, the term "mutation" carries its traditional connotation and refers to a change, inherited, naturally occurring, or introduced, in a nucleic acid or polypeptide sequence, and is used in its sense as generally known to those of skill in the art.

The term "naturally occurring", as applied to an object, refers to the fact that an object can be found in nature. For example, a polypeptide or polynucleotide sequence that is present in an organism (including bacteria) that can be isolated from a source in nature and which has not been intentionally modified by man in the laboratory is naturally occurring.

The term "nucleic acid" refers to a polymeric form of nucleotides, either ribonucleotides or deoxynucleotides or a modified form of either type of nucleotide. The terms should also be understood to include, as equivalents, analogs of either RNA or DNA made from nucleotide analogs, and, as

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applicable to the embodiment being described, single-stranded (such as sense or antisense) and double-stranded polynucleotides.

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The term "nucleic acid of the invention" refers to a nucleic acid encoding a polypeptide of the invention, i.e., a nucleic acid comprising a sequence consisting of, or consisting essentially of, the polynucleotide sequence set forth in SEQ ID NO: 1 or SEQ ID NO: 3. A nucleic acid of the invention can comprise all, or a portion of: the nucleotide sequence of SEQ ID NO: 1 or SEQ ID NO: 3; a nucleotide sequence at least 60%, 70%, 80%, 90%, 95%, 96%, 97%, 98% or 99% identical to SEQ ID NO: 1 or SEQ ID NO: 3; a nucleotide sequence that hybridizes under stringent conditions to SEQ ID NO: 1 or SEQ ID NO: 3; nucleotide sequences encoding polypeptides that are functionally equivalent to polypeptides of the invention; nucleotide sequences encoding polypeptides at least about 60%, 70%, 80%, 85%, 90%, 95%, 98%, 99% homologous or identical with an amino acid sequence of SEQ ID NO: 2 or SEQ ID NO: 4; nucleotide sequences encoding polypeptides having an activity of a polypeptide of the invention and having at least about 60%, 70%, 80%, 85%, 90%, 95%, 98%, 99% or more homology or identity with SEQ ID NO: 2 or SEQ ID NO: 4; nucleotide sequences that differ by 1 to about 2, 3, 5, 7, 10, 15, 20, 30, 50, 75 or more nucleotide substitutions, additions or deletions, such as allelic variants, of SEQ ID NO: 1 and SEQ ID NO: 3; nucleic acids derived from and evolutionarily related to SEQ ID NO: 1 or SEQ ID NO: 3; and complements of and nucleotide sequences resulting from the degeneracy of the genetic code, for all of the foregoing and other nucleic acids of the invention. Nucleic acids of the invention also include homologs. i.e., orthologs and paralogs, of SEQ ID NO: 1 or SEQ ID NO: 3 and also variants of SEQ ID NO: 1 or SEQ ID NO: 3 which have been codon optimized for expression in a particular organism (i.e., host cell).

The term "operably linked", when describing the relationship between two nucleic acid regions, refers to a juxtaposition wherein the regions are in a relationship permitting them to function in their intended manner. For example, a control sequence "operably linked" to a coding sequence is ligated in such a way that expression of the coding sequence is achieved under

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conditions compatible with the control sequences, such as when the appropriate molecules (i.e., inducers and polymerases) are bound to the control or regulatory sequence(s).

As used herein, "orthorhombic unit cell" refers to a unit cell wherein a  $\neq$  b  $\neq$  c, and  $\alpha = \beta = \gamma = 90^{\circ}$ . The vectors a, b, and c describe the unit cell edges and the angles  $\alpha$ ,  $\beta$ , and  $\gamma$  describe the unit cell angles.

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As used herein, the term "CAR" refers to any polypeptide with an amino acid sequence that can be aligned with at least one of human, mouse, or rat CAR, such that at least 50% of the amino acids are identical to the corresponding amino acid in the human, mouse, or rat CAR. The term "CAR" also encompasses nucleic acids for which the corresponding translated protein sequence can be considered to be a CAR. The term "CAR" includes vertebrate homologs of CAR family members including, but not limited to mammalian and avian homologs. Representative mammalian homologs of CAR family members include, but are not limited to murine and human homologs.

As used herein, the terms "CAR gene" and "recombinant CAR gene" are used interchangeably and refer to a nucleic acid molecule comprising an open reading frame encoding a CAR polypeptide, including both exon and (optionally) intron sequences.

As used herein, the terms "CAR gene product", "CAR protein", "CAR polypeptide", and "CAR peptide" are used interchangeably and refer to peptides having amino acid sequences which are substantially identical to native CAR amino acid sequences from the organism of interest and which are biologically active in that they comprise all or a part of the amino acid sequence of a CAR polypeptide, or cross-react with antibodies raised against a CAR polypeptide, or retain all or some of the biological activity (e.g., DNA or ligand-binding ability and/or dimerization ability) of the native amino acid sequence or protein. Such biological activity can include immunogenicity.

As used herein, the terms "CAR gene product", "CAR protein", "CAR polypeptide", and "CAR peptide" are used interchangeably and refer to a subtype of the CAR family. In one embodiment, a CAR gene product is CAR.

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In another embodiment, a CAR gene product comprises the amino acid sequence of SEQ ID NO: 2.

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As used herein, the terms "CAR gene product", "CAR protein", "CAR polypeptide", and "CAR peptide" also include analogs of a CAR polypeptide. By "analog" is intended that a DNA or peptide sequence can contain alterations relative to the sequences disclosed herein, yet retain all or some of the biological activity of those sequences. Analogs can be derived from genomic nucleotide sequences as are disclosed herein or those from other organisms, or can be created synthetically. Those skilled in the art will appreciate that other analogs, as yet undisclosed or undiscovered, can be used to design and/or construct CAR analogs. There is no need for a "CAR gene product", "CAR protein", "CAR polypeptide", or "CAR peptide" to comprise all or substantially all of the amino acid sequence of a CAR polypeptide gene product. Shorter or longer sequences are anticipated to be of use in the invention; shorter sequences are herein referred to as "segments". Thus, the terms "CAR gene product", "CAR protein", "CAR polypeptide", and "CAR peptide" also include fusion or recombinant CAR polypeptides and proteins comprising sequences of the present invention. Methods of preparing such proteins are disclosed herein and are known in the art.

The term "phenotype" refers to the entire physical, biochemical, and physiological makeup of a cell, *i.e.*, having any one trait or any group of traits.

As used herein, the term "polypeptide" refers to any polymer comprising any of the 20 protein amino acids, regardless of its size. Although "protein" is often used in reference to relatively large polypeptides and "peptide" is often used in reference to small polypeptides, usage of these terms in the art overlaps and varies. The term "polypeptide" as used herein refers to peptides, polypeptides, and proteins, unless otherwise noted. As used herein, the terms "protein", "polypeptide" and "peptide" are used interchangeably herein when referring to a gene product. The term "polypeptide", and the terms "protein" and "peptide" which are used interchangeably herein, refers to a polymer of amino acids. Exemplary

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polypeptides include gene products, naturally occurring proteins, homologs, orthologs, paralogs, fragments, as well as other equivalents, variants, and analogs of the foregoing.

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The terms "polypeptide fragment" or "fragment", when used to refer to a reference polypeptide, refers to a polypeptide in which amino acid residues are deleted as compared to the reference polypeptide itself, but where the remaining amino acid sequence is usually identical to the corresponding positions in the reference polypeptide. Such deletions can occur at the amino-terminus or carboxy-terminus of the reference polypeptide, or alternatively both. Fragments typically are at least 5, 6, 8 or 10 amino acids long, at least 14 amino acids long, at least 20, 30, 40 or 50 amino acids long, at least 75 amino acids long, or at least 100, 150, 200, 300, 500 or more amino acids long. A fragment can retain one or more of the biological activities of the reference polypeptide. In certain embodiments, a fragment can comprise a druggable region, and optionally additional amino acids on one or both sides of the druggable region, which additional amino acids can number from 5, 10, 15, 20, 30, 40, 50, or up to 100 or more residues. Further, fragments can include a sub-fragment of a specific region, which subfragment retains a function of the region from which it is derived. In one embodiment, a fragment can have immunogenic properties.

The term "polypeptide of the invention" refers to a polypeptide comprising the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4, or an equivalent or fragment thereof: *i.e.*, a polypeptide comprising a sequence consisting of, or consisting essentially of, the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4. Polypeptides of the invention include polypeptides comprising all or a portion of the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4; the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4 with 1 to about 2, 3, 5, 7, 10, 15, 20, 30, 50, 75 or more conservative amino acid substitutions; an amino acid sequence that is at least 60%, 70%, 80%, 90%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO: 2 or SEQ ID NO: 4; and functional fragments

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thereof. Polypeptides of the invention also include homologs, *i.e.*, orthologs and paralogs, of SEQ ID NO: 2 or SEQ ID NO: 4.

As used herein, the term "primer" refers to a nucleic acid comprising in one embodiment 2 or more deoxyribonucleotides or ribonucleotides, in another embodiment more than 3, in another embodiment more than 8, and in yet another embodiment at least about 20 nucleotides of an exonic or intronic region. In one embodiment, an oligonucleotide is between 10 and 30 bases in length.

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The term "purified" refers to an object species that is the predominant species present (i.e., on a molar basis it is more abundant than any other individual species in the composition). A "purified fraction" is a composition wherein the object species comprises at least about 50 percent (on a molar basis) of all species present. In making the determination of the purity of a species in solution or dispersion, the solvent or matrix in which the species is dissolved or dispersed is usually not included in such determination; instead, only the species (including the one of interest) dissolved or dispersed are taken into account. Generally, a purified composition will have one species that comprises more than about 80 percent of all species present in the composition, more than about 85%, 90%, 95%, 99% or more of all species The object species can be purified to essential homogeneity (contaminant species cannot be detected in the composition by conventional detection methods) wherein the composition consists essentially of a single species. A skilled artisan can purify a polypeptide of the invention using standard techniques for protein purification in light of the teachings herein. Purity of a polypeptide can be determined by a number of methods known to those of skill in the art, including for example, amino-terminal amino acid sequence analysis, gel electrophoresis, mass-spectrometry analysis and the methods described herein.

The terms "recombinant protein" and "recombinant polypeptide" refer to a polypeptide that is produced by recombinant DNA techniques. An example of such techniques includes when DNA encoding a polypeptide is inserted

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into a suitable expression vector that is in turn used to transform a host cell to produce the polypeptide encoded by the DNA.

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A "reference sequence" is a defined sequence used as a basis for a sequence comparison. A reference sequence can be a subset of a larger sequence, for example, as a segment of a full-length protein given in a sequence listing such as SEQ ID NO: 2 or SEQ ID NO: 4, or can comprise a complete protein sequence. Generally, a reference sequence is at least 200, 300 or 400 nucleotides in length, frequently at least 600 nucleotides in length, and often at least 800 nucleotides in length (or the protein equivalent if it is shorter or longer in length). Because two proteins can each (1) comprise a sequence (i.e., a portion of the complete protein sequence) that is similar between the two proteins, and (2) can further comprise a sequence that is divergent between the two proteins, sequence comparisons between two (or more) proteins are typically performed by comparing sequences of the two proteins over a "comparison window" to identify and compare local regions of sequence similarity.

A "comparison window," as used herein, refers to a conceptual segment of at least 20 contiguous amino acid positions wherein a protein sequence can be compared to a reference sequence of at least 20 contiguous amino acids and wherein the portion of the protein sequence in the comparison window can comprise additions or deletions (i.e., gaps) of 20 percent or less as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. Optimal alignment of sequences for aligning a comparison window can be conducted by the local homology algorithm of Smith & Waterman, 1981, by the homology alignment algorithm of Needleman & Wunsch, 1970, by the search for similarity method of Pearson & Lipman, 1988, by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, available from Accelrys, Inc., San Diego, California, United States of America), or by inspection, and the best alignment (i.e., resulting in the highest percentage of homology over the comparison window) generated by the various methods can be identified.

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The term "regulatory sequence" is a generic term used throughout the specification to refer to polynucleotide sequences, such as initiation signals, enhancers, regulators and promoters, that are necessary or desirable to affect the expression of coding and non-coding sequences to which they are operably linked. Exemplary regulatory sequences are described in Goeddel, 1990, and include, for example, the early and late promoters of SV40, adenovirus or cytomegalovirus immediate early promoter, the lac system, the trp system, the TAC or TRC system, T7 promoter whose expression is directed by T7 RNA polymerase, the major operator and promoter regions of phage lambda, the control regions for fd coat protein, the promoter for 3phosphoglycerate kinase or other glycolytic enzymes, the promoters of acid phosphatase, i.e., Pho5, the promoters of the yeast α-mating factors, the polyhedron promoter of the baculovirus system and other sequences known to control the expression of genes of prokaryotic or eukaryotic cells or their viruses, and various combinations thereof. The nature and use of such control sequences can differ depending upon the host organism. prokaryotes, such regulatory sequences generally include promoter, ribosomal binding site, and transcription termination sequences. The term "regulatory sequence" is intended to include, at a minimum, components whose presence can influence expression, and can also include additional components whose presence is advantageous, for example, leader sequences and fusion partner sequences. In certain embodiments, transcription of a polynucleotide sequence is under the control of a promoter sequence (or other regulatory sequence) that controls the expression of the polynucleotide in a cell-type in which expression is intended. It will also be understood that the polynucleotide can be under the control of regulatory sequences that are the same or different from those sequences which control expression of the naturally occurring form of the polynucleotide.

The term "reporter gene" refers to a nucleic acid comprising a nucleotide sequence encoding a protein that is readily detectable either by its presence or activity, including, but not limited to, luciferase, fluorescent protein (i.e., green fluorescent protein), chloramphenicol acetyl transferase, β-

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galactosidase, secreted placental alkaline phosphatase,  $\beta$ -lactamase, human growth hormone, and other secreted enzyme reporters. Generally, a reporter gene encodes a polypeptide not otherwise produced by the host cell, which is detectable by analysis of the cell(s), *i.e.*, by the direct fluorometric, radioisotopic or spectrophotometric analysis of the cell(s) and preferably without the need to kill the cells for signal analysis. In certain instances, a reporter gene encodes an enzyme, which produces a change in fluorometric properties of the host cell, which is detectable by qualitative, quantitative, or semiquantitative function or transcriptional activation. Exemplary enzymes include esterases,  $\beta$ -lactamase, phosphatases, peroxidases, proteases (tissue plasminogen activator or urokinase) and other enzymes whose function can be detected by appropriate chromogenic or fluorogenic substrates known to those skilled in the art or developed in the future.

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The term "sequence homology" refers to the proportion of base matches between two nucleic acid sequences or the proportion of amino acid matches between two amino acid sequences. When sequence homology is expressed as a percentage, i.e., 50%, the percentage denotes the proportion of matches over the length of sequence from a desired sequence (i.e., SEQ. ID NO: 1) that is compared to some other sequence. Gaps (in either of the two sequences) are permitted to maximize matching; gap lengths of 15 bases or less are usually used, 6 bases or less are used more frequently, with 2 bases or less used even more frequently. The term "sequence identity" means that sequences are identical (i.e., on a nucleotide-by-nucleotide basis for nucleic acids or amino acid-by-amino acid basis for polypeptides) over a window of comparison. The term "percentage of sequence identity" is calculated by comparing two optimally aligned sequences over the comparison window, determining the number of positions at which the identical amino acids occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the comparison window, and multiplying the result by 100 to yield the percentage of sequence identity. Methods to calculate

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sequence identity are known to those of skill in the art and described in further detail herein.

As used herein, the term "sequencing" refers to determining the ordered linear sequence of nucleotides or amino acids of a DNA, RNA, or protein target sample, using conventional manual or automated laboratory techniques.

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The term "small molecule" refers to a compound, which has a molecular weight of less than about 5 kilodalton (kD), less than about 2.5 kD, less than about 1.5 kD, or less than about 0.9 kD. Small molecules can be, for example, nucleic acids, peptides, polypeptides, peptide nucleic acids, peptidomimetics, carbohydrates, lipids, or other organic (carbon containing) or inorganic molecules. The term "small organic molecule" refers to a small molecule that is often identified as being an organic or medicinal compound, and does not include molecules that are exclusively nucleic acids, peptides, or polypeptides.

The term "soluble" as used herein with reference to a polypeptide of the invention or other protein means that upon expression in cell culture, at least some portion of the polypeptide or protein expressed remains in the cytoplasmic fraction of the cell and does not fractionate with the cellular debris upon lysis and centrifugation of the lysate. Solubility of a polypeptide can be increased by a variety of art recognized methods, including fusion to a heterologous amino acid sequence, deletion of amino acid residues, amino acid substitution (i.e., enriching the sequence with amino acid residues having hydrophilic side chains), and chemical modification (i.e., addition of hydrophilic groups). The solubility of polypeptides can be measured using a variety of art recognized techniques, including dynamic light scattering to determine aggregation state. UV absorption, centrifugation to separate aggregated from non-aggregated material, and SDS gel electrophoresis (i.e., the amount of protein in the soluble fraction is compared to the amount of protein in the soluble and insoluble fractions combined). When expressed in a host cell, the polypeptides of the invention can be at least about 1%, 2%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90% or more soluble, i.e.,

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at least about 1%, 2%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90% or more of the total amount of protein expressed in the cell is found in the cytoplasmic fraction. In certain embodiments, a one liter culture of cells expressing a polypeptide of the invention will produce at least about 0.1, 0.2, 0.5, 1, 2, 5, 10, 20, 30, 40, 50 milligrams or more of soluble protein. In an exemplary embodiment, a polypeptide of the invention is at least about 10% soluble and will produce at least about 1 milligram of protein from a one liter cell culture.

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As used herein, the term "space group" refers to the arrangement of symmetry elements of a crystal.

The term "specifically hybridizes" refers to detectable and specific nucleic acid binding. Polynucleotides, oligonucleotides, and nucleic acids of the invention selectively hybridize to nucleic acid strands under hybridization and wash conditions that minimize appreciable amounts of detectable binding to nonspecific nucleic acids. Stringent conditions can be used to achieve selective hybridization conditions as known in the art and discussed herein. Generally, the nucleic acid sequence homology between the polynucleotides, oligonucleotides, and nucleic acids of the invention and a nucleic acid sequence of interest will be at least 30%, 40%, 50%, 60%, 70%, 80%, 85%, 90%, 95%, 98%, 99%, or more. In certain instances, hybridization and washing conditions are performed under stringent conditions according to conventional hybridization procedures and as described further herein.

As used herein, the terms "structure coordinates", "atomic coordinates", and "structural coordinates" are used interchangeably and refer to coordinates derived from mathematical equations related to the patterns obtained on diffraction of a monochromatic beam of X-rays by the atoms (scattering centers) of a molecule in crystal form. The diffraction data are used to calculate an electron density map of the repeating unit of the crystal. The electron density maps are used to establish the positions of the individual atoms within the unit cell of the crystal.

Those of skill in the art understand that a set of coordinates determined by X-ray crystallography is not without experimental error. In general, the

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error in the coordinates tends to be reduced as the resolution is increased, since more experimental diffraction data is available for the model fitting and refinement. Thus, for example, more diffraction data can be collected from a crystal that diffracts to a resolution of 2.0 angstroms than from a crystal that diffracts to a lower resolution, such as 2.5 or 3.0 angstroms. Consequently, the refined structural coordinates will usually be more accurate when fitted and refined using data from a crystal that diffracts to higher resolution. The design of ligands for a CAR polypeptide depends on the accuracy of the structural coordinates. If the coordinates are not sufficiently accurate, then the design process will be ineffective. In most cases, it is very difficult or impossible to collect sufficient diffraction data to define atomic coordinates precisely when the crystals diffract to a resolution of 3.0 angstroms or poorer. Thus, in most cases, it is difficult to use X-ray structures in structure-based ligand design when the X-ray structures are based on crystals that diffract to a resolution of only 3.0 angstroms or poorer. However, common experience has shown that crystals diffracting to 2.0-2.5 angstroms or better can yield Xray structures with sufficient accuracy to greatly facilitate structure-based drug design. Further improvement in the resolution can further facilitate structurebased design, but the coordinates obtained at 2.0-2.5 angstroms resolution are generally considered adequate for most purposes.

Also, those of skill in the art will understand that nuclear receptors can adopt different conformations when different ligands are bound, or in the absence of any ligand. In particular, in most nuclear receptors, the AF2 helix can adopt different conformations when agonists and antagonists (or inverse agonists) are bound. More subtle conformational changes occur in other parts of the LBD when the AF2 helix is shifted. Generally, structure-based design of ligands that modulate CAR activity requires an understanding of the "activated" conformation that occurs when agonists are bound (or in the absence of ligand), as well as the "repressed" conformation that occurs when antagonists (or inverse agonists) are bound. The crystal structure of CAR bound to Compound 1 provides the "repressed" structure of CAR. In one embodiment, the "activated" conformation of CAR can be modeled

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approximately by using the "repressed" CAR structure as a starting structure, and then adjusting the conformation of the residues at the C-terminal end of the structure, residues 332-348, to form an AF2 helix with conformation, position, and orientation similar to that observed in the "activated" conformations of other nuclear receptors. It should be noted that the X-ray structure of CAR bound to Compound 1, which is an inverse agonist, revealed a completely novel, unexpected conformation for the residues that normally comprise the AF2 helix and the AF2 linking segment. No conventional modeling procedure could have predicted this novel "repressed" structure from an X-ray structure of the "activated" conformation of CAR.

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The terms "stringent conditions" or "stringent hybridization conditions" refer to conditions that promote specific hybridization between two complementary polynucleotide strands so as to form a duplex. Stringent conditions can be selected to be about 5°C lower than the thermal melting point (Tm) for a given polynucleotide duplex at a defined ionic strength and pH. The length of the complementary polynucleotide strands and their GC content will determine the Tm of the duplex, and thus the hybridization conditions necessary for obtaining a desired specificity of hybridization. The Tm is the temperature (under defined ionic strength and pH) at which 50% of a polynucleotide sequence hybridizes to a perfectly matched complementary strand. In certain cases it can be desirable to increase the stringency of the hybridization conditions to be about equal to the Tm for a particular duplex.

A variety of techniques for estimating the Tm are available. Typically, G-C base pairs in a duplex are estimated to contribute about 3°C to the Tm, while A-T base pairs are estimated to contribute about 2°C, up to a theoretical maximum of about 80-100°C. However, more sophisticated models of Tm are available in which G-C stacking interactions, solvent effects, the desired assay temperature and the like are taken into account. For example, probes can be designed to have a dissociation temperature (Td) of approximately 60°C, using the formula: Td = ((((((3 x #GC) + (2 x #AT)) x 37) - 562)/#bp) - 5; where #GC, #AT, and #bp are the number of guanine-cytosine base pairs, the

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number of adenine-thymine base pairs, and the number of total base pairs, respectively, involved in the formation of the duplex.

Hybridization can be carried out in 5x SSC, 4x SSC, 3x SSC, 2x SSC, 1x SSC or 0.2x SSC for at least about 1 hour, 2 hours, 5 hours, 12 hours, or 24 hours. The temperature of the hybridization can be increased to adjust the stringency of the reaction, for example, from about 25°C (room temperature), to about 45°C, 50°C, 55°C, 60°C, or 65°C. The hybridization reaction can also include another agent affecting the stringency; for example, hybridization conducted in the presence of 50% formamide increases the stringency of hybridization at a defined temperature.

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The hybridization reaction can be followed by a single wash step, or two or more wash steps, which can be at the same or a different salinity and temperature. For example, the temperature of the wash can be increased to adjust the stringency from about 25°C (room temperature), to about 45°C, 50°C, 55°C, 60°C, 65°C, or higher. The wash step can be conducted in the presence of a detergent, *i.e.*, 0.1 or 0.2% SDS. For example, hybridization can be followed by two wash steps at 65°C each for about 20 minutes in 2x SSC, 0.1% SDS, and optionally two additional wash steps at 65°C each for about 20 minutes in 0.2x SSC, 0.1% SDS.

Exemplary stringent hybridization conditions include overnight hybridization at 65°C in a solution comprising, or consisting of, 50% formamide, 10x Denhardt's Solution (0.2% Ficoll, 0.2% Polyvinylpyrrolidone, 0.2% bovine serum albumin) and 200 µg/ml of denatured carrier DNA, *i.e.*, sheared salmon sperm DNA, followed by two wash steps at 65°C each for about 20 minutes in 2x SSC, 0.1% SDS, and two wash steps at 65°C each for about 20 minutes in 0.2x SSC, 0.1% SDS.

Hybridization can include hybridizing two nucleic acids in solution, or a nucleic acid in solution to a nucleic acid attached to a solid support, *i.e.*, a filter. When one nucleic acid is on a solid support, a prehybridization step can be conducted prior to hybridization. Prehybridization can be carried out for at least about 1 hour, 3 hours or 10 hours in the same solution and at the same

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temperature as the hybridization solution (without the complementary polynucleotide strand).

Appropriate stringency conditions are known to those skilled in the art or can be determined experimentally by the skilled artisan. See e.g. Ausubel et al., 1994; Sambrook & Russell, 2001; Agrawal, 1993; Tibanyenda et al., 1984; Ebel et al., 1992.

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The term "structural motif", when used in reference to a polypeptide, refers to a polypeptide that, although it can have different amino acid sequences, can result in a similar structure, wherein by structure is meant that the motif forms generally the same tertiary structure, or that certain amino acid residues within the motif, or alternatively their backbone or side chains (which can or can not include the  $C\alpha$  atoms of the side chains) are positioned in a like relationship with respect to one another in the motif.

As applied to proteins, the term "substantial identity" means that two protein sequences, when optimally aligned, such as by the programs GAP or BESTFIT using default gap weights, typically share at least about 70 percent sequence identity, alternatively at least about 80, 85, 90, 95 percent sequence identity or more. In certain instances, residue positions that are not identical differ by conservative amino acid substitutions, which are described above.

As used herein, the term "substantially pure" refers to a polynucleotide or polypeptide that is substantially free of the sequences and molecules with which it is associated in its natural state, as well as from those molecules used in the isolation procedure. The term "substantially free" refers to that the sample is in one embodiment at least 50%, in another embodiment at least 70%, in another embodiment at least 80%, and in still another embodiment at least 90% free of the sequences and molecules with which is it associated in nature.

As used herein, the term "target cell" refers to a cell, into which it is desired to insert a nucleic acid sequence or polypeptide, or to otherwise effect a modification from conditions known to be present in the unmodified cell. A nucleic acid sequence introduced into a target cell can be of variable length.

Additionally, a nucleic acid sequence can enter a target cell as a component of a plasmid or other vector or as a naked sequence.

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The term "test compound" refers to a molecule to be tested by one or more screening method(s) as a putative modulator of a polypeptide of the invention or other biological entity or process. A test compound is usually not known to bind to a target of interest. The term "control test compound" refers to a compound known to bind to the target (i.e., a known agonist, antagonist, partial agonist or inverse agonist). The term "test compound" does not include a chemical added as a control condition that alters the function of the target to determine signal specificity in an assay. Such control chemicals or conditions include chemicals that 1) nonspecifically or substantially disrupt protein structure (i.e., denaturing agents (i.e., urea or guanidinium), chaotropic agents, sulfhydryl reagents (i.e., dithiothreitol and β-mercaptoethanol), and proteases), 2) generally inhibit cell metabolism (i.e., mitochondrial uncouplers) and 3) non-specifically disrupt electrostatic or hydrophobic interactions of a protein (i.e., high salt concentrations, or detergents at concentrations sufficient to non-specifically disrupt hydrophobic interactions). Further, the term "test compound" also does not include compounds known to be unsuitable for a therapeutic use for a particular indication due to toxicity of the subject. In certain embodiments, various predetermined concentrations of test compounds are used for screening such as 0.01 μM, 0.1 μM, 1.0 μM, and Examples of test compounds include, but are not limited to peptides, nucleic acids, carbohydrates, and small molecules. The term "novel test compound" refers to a test compound that is not in existence as of the filing date of this application. In certain assays using novel test compounds, the novel test compounds comprise at least about 50%, 75%, 85%, 90%, 95% or more of the test compounds used in the assay or in any particular trial of the assay.

The term "therapeutically effective amount" refers to that amount of a modulator, drug, or other molecule that is sufficient to effect treatment when administered to a subject in need of such treatment. The therapeutically effective amount will vary depending upon the subject and disease condition

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being treated, the weight and age of the subject, the severity of the disease condition, the manner of administration and the like, which can readily be determined by one of ordinary skill in the art.

The term "transfection" means the introduction of a nucleic acid, *i.e.*, an expression vector, into a recipient cell, which in certain instances involves nucleic acid-mediated gene transfer. The term "transformation" refers to a process in which a cell's genotype is changed as a result of the cellular uptake of exogenous nucleic acid. For example, a transformed cell can express a recombinant form of a polypeptide of the invention or antisense expression can occur from the transferred gene so that the expression of a naturally occurring form of the gene is disrupted.

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The term "transgene" means a nucleic acid sequence, which is partly or entirely heterologous to a transgenic animal or cell into which it is introduced, or, is homologous to an endogenous gene of the transgenic animal or cell into which it is introduced, but which is designed to be inserted, or is inserted, into the animal's genome in such a way as to alter the genome of the cell into which it is inserted (i.e., it is inserted at a location which differs from that of the natural gene or its insertion results in a knockout). A transgene can include one or more regulatory sequences and any other nucleic acids, such as introns, that can be necessary for optimal expression.

The term "transgenic animal" refers to any animal, for example, a mouse, rat or other non-human mammal, a bird or an amphibian, in which one or more of the cells of the animal contain heterologous nucleic acid introduced by way of human intervention, such as by transgenic techniques well known in the art. The nucleic acid is introduced into the cell, directly or indirectly, by way of deliberate genetic manipulation, such as by microinjection or by infection with a recombinant virus. The term genetic manipulation does not include classical cross-breeding, or *in vitro* fertilization, but rather is directed to the introduction of a recombinant DNA molecule. This molecule can be integrated within a chromosome, or it can be extrachromosomally replicating DNA. In the typical transgenic animals described herein, the transgene

causes cells to express a recombinant form of a protein. However, transgenic animals in which the recombinant gene is silent are also contemplated.

As used herein, the term "unit cell" refers to a basic parallelepiped shaped block. Each unit cell comprises a complete representation of the unit of pattern, the repetition of which builds up the crystal. Thus, the term "unit cell" refers to the fundamental portion of a crystal structure that is repeated infinitely by translation in three dimensions. A unit cell is characterized by three vectors, a, b, and c, not located in one plane, which form the edges of a parallelepiped. Angles  $\alpha$ ,  $\beta$  and  $\gamma$  define the angles between the vectors: angle  $\alpha$  is the angle between vectors b and c; angle  $\beta$  is the angle between vectors a and c; and angle  $\gamma$  is the angle between vectors a and b. The entire volume of a crystal can be constructed by regular assembly of unit cells, each unit cell comprising a complete representation of the unit of pattern, the repetition of which builds up the crystal.

Unless otherwise indicated, all numbers expressing quantities of ingredients, reaction conditions, and so forth used in the specification and claims are to be understood as being modified in all instances by the term "about". Accordingly, unless indicated to the contrary, the numerical parameters set forth in this specification and attached claims are approximations that can vary depending upon the desired properties sought to be obtained by the present invention.

### II. Description of Tables

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Table 1 is a table summarizing the crystal and data statistics obtained from the crystallized ligand-binding domain of CAR in complex with the ligand Compound 1. Data on the unit cell are presented, including data on the crystal space group, unit cell dimensions, molecules per asymmetric cell and crystal resolution.

Table 2 is a table of the atomic coordinate data obtained from X-ray diffraction from the ligand-binding domain of CAR in complex with the ligand Compound 1.

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Table 3 is a table of the atomic structure coordinate data of the polyalanine model of the conserved vitamin D receptor ligand-binding domain.

### III. General Considerations

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The present invention is applicable *mutatis mutandis* to all CARs, as discussed herein, based in part on the patterns of CAR structure and modulation that have emerged as a consequence of determining the three dimensional structure of CAR with bound ligand. Analysis and alignment of amino acid sequences, and X-ray and NMR structure determinations, have shown that nuclear receptors have a modular architecture with three main domains:

- 1) a variable amino-terminal domain;
- 2) a highly conserved DNA-binding domain (DBD); and
- 3) a less conserved carboxy-terminal ligand-binding domain (LBD).

In addition, nuclear receptors can have linker segments of variable length between these major domains. Sequence analysis and X-ray crystallography, including the work of the present invention, have confirmed that CARs, and indeed many NRs, also have the same general modular architecture, with the same three domains. The function of the CARs in human cells presumably requires all three domains in a single amino acid sequence. However, the modularity of the CARs permits different domains of each protein to separately accomplish certain functions.

Previous analysis of the nuclear receptors has revealed multiple discrete functional modules within the family that display generalized functional characteristics (for review see Beato et al., 1995; Kastner et al., 1995; Mangelsdorf & Evans, 1995; Tzukerman et al., 1994). A variable amino-terminal domain (A/B) is present that sometimes contains a strong and autonomous activation function (AF1), shown to be critical for cell and target gene specificity (Tora et al., 1988). A more carboxyl-terminal central region contains a DNA binding domain (DBD) characterized by two C4-type zinc fingers. The DBD binds to specific genomic response elements and thereby regulates the transcriptional activity of select genes containing the response

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elements. At the distal carboxyl terminus, a ligand-binding domain (LBD) is present containing a highly conserved second transactivation function (AF2) that is important for hormone-dependent transcriptional transactivation (Lanz & Rusconi, 1994).

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Typically, the LBD forms a three-layered anti-parallel helical sandwich composed of 10-14 α helices and a β-sheet with 2-4 strands. The helices pack together so as to leave a binding pocket near the middle of the bundle, capped on one side by the β-sheet, and, in the "activated" state, capped on the other side by the AF2-helix. Comparison of apo, agonist-bound, and antagonist-bound nuclear receptor structures has led to a model for ligandinducible receptor action. In this model, the agonist (activating) ligands tend to hold the AF2 helix in a conformation where it "caps" the binding pocket. Antagonistic ligands usually shift the AF2 helix out of this "active" position. The AF2 helix can also shift into other conformations, positions, and orientations in the absence of ligand. Constitutively active receptors such as CAR should presumably utilize a similar mechanism of action, except that the AF2 helix adopts the "active" position, capping the ligand-binding pocket, even in the absence of ligand. Inverse agonists would presumably tend to shift the AF2 helix out of this "active" position, whereas superagonists would presumably tend to hold the AF2 helix more tightly in the active position. Central to the efficient ligand-induced transcriptional activation is the recruitment of co-regulator proteins - coactivators and co-repressors, which interact with the LBD and activate or repress transactivation, respectively (Moras & Gronemeyer, 1998; Weatherman et al., 1999; McKenna & O'Malley, 2000). In general, the conformational changes described above involving the AF2 helix cause changes in the affinity of the LBD for co-repressors versus coactivators. The binding of an agonist results in a dissociation of corepressors and brings the AF2 into a context where it can interact with transcriptional coactivators. Likewise, an antagonist would be expected to disrupt the binding of coactivators.

Sequences that function in nuclear localization, receptor dimerization, and interaction with heat-shock proteins (Gronemeyer & Laudet, 1995) are

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also present within the nuclear receptor substructure. Through the coordinated action of these separate functional domains, nuclear receptor activation by ligand culminates in modulation of target gene expression through DNA interactions (Tsai & O'Malley, 1994) or in certain other cases through cross-talk with other cell signaling pathways (Stein & Yang, 1995; Paech et al., 1998). In short, a ligand alters nuclear receptor function by altering the conformation of the receptor and consequently the constellation of protein-protein interactions in which the receptor is engaged (Freedman, 1999).

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Some of the functions of a domain within the full-length receptor are preserved when that particular domain is isolated from the remainder of the protein. Using conventional protein chemistry techniques, a modular domain can sometimes be separated from the parent protein. Using conventional molecular biology techniques, each domain can usually be separately expressed with its original function intact or, as discussed herein below, chimeras comprising two different proteins can be constructed, wherein the chimeras retain the properties of the individual functional domains of the respective nuclear receptors from which the chimeras were generated.

The LBD is the second most highly conserved domain in these 3 domains. As its name suggests, the LBD binds ligands. With many nuclear receptors binding of the ligand can induce a conformational change in the LBD that can, in turn, increase or decrease transcription of certain target genes. The LBD also participates in other functions, including dimerization and nuclear translocation.

X-ray structures have shown that most nuclear receptor LBDs adopt the same general folding pattern. This fold includes 10-12 alpha helices arranged in a bundle, together with several beta-strands, additional alpha helices and linking segments. The major alpha helices and beta-strands have been numbered differently in different publications. The present disclosure follows the numbering scheme of Nolte *et al.*, 1998, where the major alphahelices and beta-strands in PPARγ were designated sequentially through the amino acid sequence as H1, H2, S1, H2', H3, H3', H4, H5, S2, S3, S4, H6,

H7, H8, H9, H10 and HAF. The alpha helix at the C-terminal end, HAF, is also called "helix-AF", "helix-AF2" the "AF2 helix" or "helix-12". Most, but not all, of these alpha helices and beta-strands are observed in the structure of CAR. An additional helix, designated here as "helix-X", is observed in the structure of CAR bound to Compound 1 on the C-terminal side of H10.

As described herein, the LBD of a CAR can be expressed, crystallized, its three dimensional structure determined with a ligand bound as disclosed in the present invention, and computational methods can be used to design ligands to its LBD.

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# IV. Synthesis of CAR Ligands and Intermediates

# IV.A. Compound 1 – An Embodiment of a Synthetic CAR Ligand

In one embodiment, the present invention provides compounds of Compound 1 (Formula (A) below) and tautomeric forms, pharmaceutically acceptable salts and solvates thereof:

# IV.B. Synthesis of Compound 1 and Intermediates

Compound 1, which was co-crystallized with the CAR LBD in the present invention, can be prepared as described in Example 6 and shown in

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Figure 7. Briefly, a solution of 3-fluoro-4-nitrobenzoic acid in anhydrous N.Ndimethylformamide was treated with [O-(7-azabenzotriazol-1-yl)-1,1,3,3tetramethyluronium hexafluorophosphate] followed by N.Ndiisopropylethylamine. After shaking for 5 minutes, the mixture was added to polystyrene Rink amide AM resin, and the reaction was rotated at 25°C for 18 hours. The reaction solution was drained, and the resin was washed with N,N-dimethylformamide, dichloromethane, methanol, and dichloromethane. The dried resin was treated with a 0.5 M phenethylamine in Nmethylpyrrolidinone solution and incubated with rotation for 15 hours at 70°C. The reaction was cooled to room temperature, drained, and the resin was washed as before. The resin was then treated with a 2.0 M SnCl<sub>2</sub>•dihydrate in N-methylpyrrolidinone solution for 24 hours at 25°C with rotation. The reaction was drained and the resin washed with 30% ethylenediamine, N,Ndimethylformamide, dichloromethane, methanol, and dichloromethane. The dried diamine resin was treated with a 0.5 M benzyhydryl isothiocyanate in Nmethylpyrrolidinone solution and a 1.0 M diisopropylcarbodiimide in Nmethylpyrrolidinone solution at 80°C with rotation. After 24 hours, the reaction was cooled to 25°C, drained, and the resin was washed with N,Ndimethylformamide, dichloromethane, methanol, and dichloromethane. The resin was then treated with 95:5 TFA:H<sub>2</sub>O and rotated at 25°C for 3 hours. The resin was drained and washed with dichloromethane. The filtrate was The oil was redissolved in concentrated in vacuo to give an oil. dichloromethane and the solution was washed twice with saturated sodium bicarbonate. The organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and concentrated in vacuo. The crude product was triturated with Et<sub>2</sub>O/hexanes, and the solid was collected by filtration to give Compound 1 as an off-white solid.

#### V. Production of CAR Polypeptides

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The native and mutated CAR polypeptides, and fragments thereof, of the present invention can be chemically synthesized in whole or part using techniques that are well known in the art (see e.g., Creighton, 1983, incorporated herein in its entirety). Alternatively, methods which are well

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known to those skilled in the art can be used to construct expression vectors containing a partial or the entire native or mutated CAR polypeptide coding sequence and appropriate transcriptional/translational control signals. These methods include *in vitro* recombinant DNA techniques, synthetic techniques, and *in vivo* recombination/genetic recombination (see e.g., the techniques described throughout Sambrook & Russell, 2001, and Ausubel *et al.*, 1994, both incorporated herein in their entirety).

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A variety of host-expression vector systems can be utilized to express a CAR coding sequence. These include but are not limited to microorganisms such as bacteria transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA expression vectors containing a CAR coding sequence; yeast transformed with recombinant yeast expression vectors containing a CAR coding sequence; insect cell systems infected with recombinant virus expression vectors (e.g., baculovirus) containing a CAR coding sequence; plant cell systems infected with recombinant virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or transformed with recombinant plasmid expression vectors (e.g., Ti plasmid) containing a CAR coding sequence; or animal cell systems. The expression elements of these systems vary in their strength and specificities.

Depending on the host/vector system utilized, any of a number of suitable transcription and translation elements, including constitutive and inducible promoters, can be used in the expression vector. For example, when cloning in bacterial systems, inducible promoters such as pL of bacteriophage  $\lambda$ , plac, ptrp, ptac (ptrp-lac hybrid promoter) and the like can be used. When cloning in insect cell systems, promoters such as the baculovirus polyhedrin promoter can be used. When cloning in plant cell systems, promoters derived from the genome of plant cells, such as heat shock promoters; the promoter for the small subunit of ribulose bisphosphate carboxylase (RUBISCO); the promoter for the chlorophyll a/b binding protein; or from plant viruses (e.g., the 35S RNA promoter of CaMV; the coat protein promoter of TMV) can be used. When cloning in mammalian cell systems, promoters derived from the genome of mammalian cells (e.g., metallothionein

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promoter) or from mammalian viruses (e.g., the adenovirus late promoter; the vaccinia virus 7.5K promoter) can be used.

In each of these systems, one of ordinary skill in the art will appreciate that other promoters can be used, and as such, the list presented is not intended to be exhaustive.

## VI. Analysis of Protein Properties

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VI.A. Analysis of Proteins by X-ray Crystallography Generally

## VI.A.1. X-ray Structure Determination

Exemplary methods for obtaining the three dimensional structure of the crystalline form of a molecule or complex are described herein and, in view of this specification, variations on these methods will be apparent to those skilled in the art (see Ducruix & Geige, 1992).

A variety of methods involving X-ray crystallography are contemplated by the present invention. For example, the present invention contemplates producing a crystallized polypeptide of the invention, or a fragment thereof, by: (a) introducing into a host cell an expression vector comprising a nucleic acid encoding for a polypeptide of the invention, or a fragment thereof; (b) culturing the host cell in a cell culture medium to express the polypeptide or fragment; (c) isolating the polypeptide or fragment from the cell culture; and (d) crystallizing the polypeptide or fragment thereof. Alternatively, the present invention contemplates determining the three dimensional structure of a crystallized polypeptide of the invention, or a fragment thereof, by: (a) crystallizing a polypeptide of the invention, or a fragment thereof, such that the crystals will diffract X-rays to a resolution of 2.5 Å or better; and (b) analyzing the polypeptide or fragment by X-ray diffraction to determine the three-dimensional structure of the crystallized polypeptide.

X-ray crystallography techniques generally require that the protein molecules be available in the form of a crystal. Crystals can be grown from a solution containing a purified polypeptide of the invention, or a fragment thereof (i.e., a ligand-binding domain), by a variety of conventional processes. These processes include, for example, batch, liquid, bridge, dialysis, and

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vapor diffusion (i.e., hanging drop or sitting drop methods). See e.g., McPherson, 1982; McPherson, 1990; Webe, 1991.

In certain embodiments, native crystals of the invention can be grown by adding precipitants to the concentrated solution of the polypeptide. The precipitants are added at a concentration just below that necessary to precipitate the protein. Water can be removed by controlled evaporation to produce precipitating conditions, which are maintained until crystal growth ceases.

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The formation of crystals is dependent on a number of different parameters, including pH, temperature, protein concentration, the nature of the solvent and precipitant, as well as the presence of added ions or ligands to the protein. In addition, the sequence of the polypeptide being crystallized will have a significant affect on the success of obtaining crystals. Many routine crystallization experiments can be needed to screen all these parameters for the few combinations that might give crystal suitable for X-ray diffraction analysis. See e.g., Jancarik & Kim, 1991.

Crystallization robots can automate and speed up the work of reproducibly setting up large number of crystallization experiments. Once some suitable set of conditions for growing the crystal are found, variations of the condition can be systematically screened in order to find the set of conditions which allows the growth of sufficiently large, single, well ordered crystals. In certain instances, a polypeptide of the invention is co-crystallized with a ligand: in one embodiment, Compound 1.

A number of methods are available to produce suitable radiation for X-ray diffraction. For example, X-ray beams can be produced by synchrotron rings where electrons (or positrons) are accelerated through an electromagnetic field while traveling at close to the speed of light. Because the admitted wavelength can also be controlled, synchrotrons can be used as a tunable X-ray source (Hendrickson, 2000). For less conventional Laue diffraction studies, polychromatic X-rays covering a broad wavelength window are used to observe many diffraction intensities simultaneously (Stoddard,

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1998). Neutrons can also be used for solving protein crystal structures (Gutberlet et al., 2001).

Before data collection commences, a protein crystal can be frozen to protect it from radiation damage. A number of different cryo-protectants can be used to assist in freezing the crystal, such as methyl pentanediol (MPD), isopropanol, ethylene glycol, glycerol, formate, citrate, mineral oil, or a low-molecular-weight polyethylene glycol (PEG). The present invention contemplates a composition comprising a polypeptide of the invention and a cryo-protectant. As an alternative to freezing the crystal, the crystal can also be used for diffraction experiments performed at temperatures above the freezing point of the solution. In these instances, the crystal can be protected from desiccation by placing it in a narrow capillary of a suitable material (generally glass or quartz) with some of the crystal growth solution included in order to maintain vapor pressure.

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X-ray diffraction results can be recorded by a number of ways known to one of skill in the art. Examples of area electronic detectors include charge coupled device detectors, multi-wire area detectors, and phosphoimager detectors (Amemiya, 1997; Westbrook & Naday, 1997; Kahn & Fourme, 1997).

A suitable system for laboratory data collection might include a Bruker AXS Proteum R system, equipped with a copper rotating anode source, Confocal MAX-FLUX<sup>TM</sup> optics and a SMART 6000 charge coupled device detector. Collection of X-ray diffraction patterns is well known to those skilled in the art (see e.g. Ducruix & Geige, 1992).

The theory behind diffraction by a crystal upon exposure to X-rays is well known. Because phase information is not directly measured in the diffraction experiment and is needed to reconstruct the electron density map, methods that can recover this missing information are required. One method of solving structures *ab initio* is the real/reciprocal space cycling technique. Suitable real/reciprocal space cycling search programs include Shake-and-Bake (Miller *et al.*, 1993; Weeks *et al.*, 1994).

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Other methods for deriving phases might also be needed. These techniques generally rely on the idea that if two or more measurements of the same reflection are made where strong, measurable, differences are attributable to the characteristics of a small subset of the atoms alone, then the contributions of other atoms can be, to a first approximation, ignored, and the positions of these atoms can be determined from the difference in scattering by one of the above techniques. Knowing the position and scattering characteristics of those atoms, one can calculate what phase the overall scattering must have had to produce the observed differences.

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One version of this technique is the isomorphous replacement technique, which requires the introduction of new, well ordered, X-ray scatterers into the crystal. These additions are usually heavy metal atoms. (so that they make a significant difference in the diffraction pattern); and if the additions do not change the structure of the molecule or of the crystal cell, the resulting crystals should be isomorphous. Isomorphous replacement experiments are usually performed by diffusing different heavy-metal metals into the channels of a pre-existing protein crystal. Growing the crystal from protein that has been soaked in the heavy atom is also possible (Petsko, 1985). Alternatively, the heavy atom can also be reactive and attached covalently to exposed amino acid side chains (such as the sulfur atom of cysteine) or it can be associated through non-covalent interactions. It is sometimes possible to replace endogenous light metals in metallo-proteins with heavier ones, i.e., zinc by mercury, or calcium by samarium (Petsko, 1985). Exemplary sources for such heavy compounds include, but are not limited to, sodium bromide, sodium selenate, trimethyl lead acetate, mercuric chloride. methyl mercury acetate, platinum tetracyanide, platinum tetrachloride, nickel chloride, and europium chloride.

A second technique for generating differences in scattering involves the phenomenon of anomalous scattering. X-rays that cause the displacement of an electron in an inner shell to a higher shell are subsequently rescattered, but there is a time lag that shows up as a phase delay. This phase delay is observed as a (generally quite small) difference in

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intensity between reflections known as Friedel mates that would be identical if no anomalous scattering were present. A second effect related to this phenomenon is that differences in the intensity of scattering of a given atom will vary in a wavelength-dependent manner, giving rise to what are known as dispersive differences. In principle, anomalous scattering occurs with all atoms, but the effect is strongest with heavy atoms, and can be maximized by using X-rays at a wavelength where the energy is equal to the difference in energy between shells. The technique therefore requires the incorporation of some heavy atom much as is needed for isomorphous replacement, although for anomalous scattering a wider variety of atoms are suitable, including lighter metal atoms (copper, zinc, iron) in metallo-proteins. One method for preparing a protein for anomalous scattering involves replacing the methionine residues in whole or in part with selenium-containing selenomethionine. Soaking with halide salts such as bromides and other non-reactive ions can also be effective (Dauter et al., 2001).

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In another process, known as multiple anomalous scattering or MAD, two to four suitable wavelengths of data are collected. (Hendrickson & Ogata, 1997). Phasing by various combinations of single and multiple isomorphous and anomalous scattering are possible too. For example, SIRAS (single isomorphous replacement with anomalous scattering) utilizes both the isomorphous and anomalous differences for one derivative to derive phases. More traditionally, several different heavy atoms are soaked into different crystals to get sufficient phase information from isomorphous differences while ignoring anomalous scattering, in the technique known as multiple isomorphous replacement (MIR) (Petsko, 1985).

Additional restraints on the phases can be derived from density modification techniques. These techniques use either generally known features of electron density distribution or known facts about that particular crystal to improve the phases. For example, because protein regions of the crystal scatter more strongly than solvent regions, solvent flattening/flipping can be used to adjust phases to make solvent density a uniform flat value (Zhang et al., 1997). If more than one molecule of the protein is present in the

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asymmetric unit, the fact that the different molecules should be virtually identical can be exploited to further reduce phase error using non-crystallographic symmetry averaging (Villieux & Read, 1997). Suitable programs for performing these processes include DM and other programs of the CCP4 suite (Collaborative Computational Project, 1994) and CNX.

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The unit cell dimensions, symmetry, vector amplitude and derived phase information can be used in a Fourier transform function to calculate the electron density in the unit cell, *i.e.*, to generate an experimental electron density map. This can be accomplished using programs of the CNX or CCP4 packages. The resolution is measured in Ångstrom (Å) units, and is closely related to how far apart two objects need to be before they can be reliably distinguished. The smaller this number is, the higher the resolution and therefore the greater the amount of detail that can be seen. In alternative embodiments, crystals of the invention diffract X-rays to a resolution of better than about 4.0, 3.5, 3.0, 2.5, 2.0, 1.5, 1.0, 0.5 Å, or better.

As used herein, the term "modeling" includes the quantitative and qualitative analysis of molecular structure and/or function based on atomic structural information and interaction models. The term "modeling" includes conventional numeric-based molecular dynamic and energy minimization models, interactive computer graphic models, modified molecular mechanics models, distance geometry and other structure-based constraint models.

Model building can be accomplished by either the crystallographer using a computer graphics program such as TURBO or O (Jones *et al.*, 1991) or, under suitable circumstances, by using a fully automated model building program, such as wARP (Perrakis *et al.*, 1999) or MAID (Levitt, 2001). This structure can be used to calculate model-derived diffraction amplitudes and phases. The model-derived and experimental diffraction amplitudes can be compared and the agreement between them can be described by a parameter referred to as R-factor. A high degree of correlation in the amplitudes corresponds to a low R-factor value, with 0.0 representing exact agreement and 0.59 representing a completely random structure. Because the R-factor can be lowered by introducing more free parameters into the model, an

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unbiased, cross-correlated version of the R-factor known as the R-free gives a more objective measure of model quality. For the calculation of this parameter a subset of reflections (generally around 10%) are set aside at the beginning of the refinement and not used as part of the refinement target. These reflections are then compared to those predicted by the model (Kleywegt & Brunger, 1996).

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The model can be improved using computer programs that maximize the probability that the observed data was produced from the predicted model. while simultaneously optimizing the model geometry. For example, the CNX program can be used for model refinement, as can the XPLOR program (Murshudov et al., 1997). In order to maximize the convergence radius of refinement, simulated annealing refinement using torsion angle dynamics can be employed in order to reduce the degrees of freedom of motion of the model (Adams et al., 1997). Where experimental phase information is available (i.e., where MAD data was collected) Hendrickson-Lattman phase probability targets can be employed. Isotropic or anisotropic domain, group or individual temperature factor refinement, can be used to model variance of the atomic position from its mean. Well-defined peaks of electron density not attributable to protein atoms are generally modeled as water molecules. Water molecules can be found by manual inspection of electron density maps, or with automatic water picking routines. Additional small molecules. including ions, cofactors, buffer molecules, or substrates can be included in the model if sufficiently unambiguous electron density is observed in a map.

In general, the R-free is rarely as low as 0.15 and can be as high as 0.35 or greater for a reasonably well-determined protein structure. The residual difference is a consequence of approximations in the model (inadequate modeling of residual structure in the solvent, modeling atoms as isotropic Gaussian spheres, assuming all molecules are identical rather than having a set of discrete conformers, etc.) and errors in the data (Lattman, 1996). In refined structures at high resolution, there are usually no major errors in the orientation of individual residues, and the estimated errors in atomic positions are usually around 0.1 - 0.2 up to 0.3 Å.

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The three dimensional structure of a new crystal can be modeled using molecular replacement. The term "molecular replacement" refers to a method that involves generating a preliminary model of a molecule or complex whose structure coordinates are unknown, by orienting and positioning a molecule whose structure coordinates are known within the unit cell of the unknown crystal, so as best to account for the observed diffraction pattern of the unknown crystal. Phases can then be calculated from this model and combined with the observed amplitudes to give an approximate Fourier synthesis of the structure whose coordinates are unknown. This, in turn, can be subject to any of the several forms of refinement to provide a final, accurate structure of the unknown crystal (Lattman, 1985; Rossmann, 1972).

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Commonly used computer software packages for molecular replacement are CNX, X-PLOR (Brunger 1992, *Nature* 355: 472-475), AMORE (Navaza, 1994, *Acta Crystallogr.* A50:157-163), the CCP4 package, the MERLOT package (Fitzgerald, 1988) and XTALVIEW (McCree *et al.*, 1992). The quality of the model can be analyzed using a program such as PROCHECK or 3D-Profiler (Laskowski *et al.*, 1993; Luthy *et al.*, 1992; Bowie *et al.*, 1991).

Homology modeling (also known as comparative modeling or knowledge-based modeling) methods can also be used to develop a three dimensional model from a polypeptide sequence based on the structures of known proteins. The method utilizes a computer model of a known protein, a computer representation of the amino acid sequence of the polypeptide with an unknown structure, and standard computer representations of the structures of amino acids. This method is well known to those skilled in the art (Greer, 1985; Blundell et al., 1988; Knighton et al., 1992). Computer programs that can be used in homology modeling are QUANTA and the Homology module in the Insight II modeling package distributed by Molecular Simulations Inc. (now part of Accelrys Inc., San Diego, California, United States of America), or MODELLER (Rockefeller University, New York, New York, United States of America). These computer programs can also be used

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for computational loop modeling techniques. See also Tosatto et al., 2002; Fiser et al., 2000.

Once a homology model has been generated it is analyzed to determine its correctness. A computer program available to assist in this analysis is the Protein Health module in QUANTA that provides a variety of tests. Other programs that provide structure analysis along with output include PROCHECK and 3D-Profiler (Luthy et al., 1992; Bowie et al., 1991). Once any irregularities have been resolved, the entire structure can be further refined.

Other molecular modeling techniques can also be employed in accordance with this invention. See e.g., Cohen et al., 1990; Navia & Murcko, 1992.

Under suitable circumstances, the entire process of solving a crystal structure can be accomplished in an automated fashion by a system such as ELVES (http://ucxray.berkeley.edu/~jamesh/elves/index.html) with little or no user intervention.

### VI.A.2. X-ray Structure

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The present invention provides methods for determining some or all of the structural coordinates for amino acids of a polypeptide of the invention, or a complex thereof.

In another aspect, the present invention provides methods for identifying a druggable region of a polypeptide of the invention. For example, one such method includes: (a) obtaining crystals of a polypeptide of the invention or a fragment thereof such that the three dimensional structure of the crystallized protein can be determined to a resolution of 2.5 Å or better; (b) determining the three dimensional structure of the crystallized polypeptide or fragment using X-ray diffraction; and (c) identifying a druggable region of a polypeptide of the invention based on the three-dimensional structure of the polypeptide or fragment.

A three dimensional structure of a molecule or complex can be described by the set of atoms that best predict the observed diffraction data

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(that is, which possesses a minimal R value). Files can be created for the structure that defines each atom by its chemical identity, spatial coordinates in three dimensions, root mean squared deviation from the mean observed position and fractional occupancy of the observed position.

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Those of skill in the art understand that a set of structure coordinates for a protein, complex, or a portion thereof, is a relative set of points that define a shape in three dimensions. Thus, it is possible that an entirely different set of coordinates could define a similar or identical shape. Moreover, slight variations in the individual coordinates can have little affect on overall shape. Such variations in coordinates can be generated because of mathematical manipulations of the structure coordinates. For example, structure coordinates could be manipulated by crystallographic permutations of the structure coordinates, fractionalization of the structure coordinates, integer additions or subtractions to sets of the structure coordinates, inversion of the structure coordinates or any combination of the above. Alternatively, modifications in the crystal structure due to mutations, additions, substitutions. and/or deletions of amino acids, or other changes in any of the components that make up the crystal, could also yield variations in structure coordinates. Such slight variations in the individual coordinates will have little affect on overall shape. If such variations are within an acceptable standard error as compared to the original coordinates, the resulting three-dimensional shape is considered to be structurally equivalent. It should be noted that slight variations in individual structure coordinates of a polypeptide of the invention or a complex thereof would not be expected to significantly alter the nature of modulators that could associate with a druggable region thereof. Thus, for example, a modulator that bound to the active site of a polypeptide of the invention would also be expected to bind to or interfere with another active site whose structure coordinates define a shape that falls within the acceptable error.

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A crystal structure of the present invention can be used to make a structural or computer model of the polypeptide, complex, or portion thereof. A model can represent the secondary, tertiary, and/or quaternary structure of

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the polypeptide, complex, or portion. The configurations of points in space derived from structure coordinates according to the invention can be visualized as, for example, a holographic image, a stereodiagram, a model, or a computer-displayed image, and the invention thus includes such images, diagrams, or models.

#### VI.A.3. Structural Equivalents

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Various computational analyses can be used to determine whether a molecule or the active site portion thereof is structurally equivalent with respect to its three-dimensional structure, to all or part of a structure of a polypeptide of the invention or a portion thereof.

For the purpose of this invention, any molecule or complex or portion thereof, that has a root mean square deviation of conserved residue backbone atoms (N, C $\alpha$ , C, O) of less than about 1.75 Å, when superimposed on the relevant backbone atoms described by the reference structure coordinates of a polypeptide of the invention, is considered "structurally equivalent" to the reference molecule. That is to say, the crystal structures of those portions of the two molecules are substantially identical, within acceptable error. Alternatively, the root mean square deviation can be is less than about 1.50, 1.40, 1.25, 1.0, 0.75, 0.5 or 0.35 Å.

The term "root mean square deviation" is understood in the art and means the square root of the arithmetic mean of the squares of the deviations. It is a way to express the deviation or variation from a trend or object.

In another aspect, the present invention provides a scalable three-dimensional configuration of points, at least a portion of said points, and preferably all of said points, derived from structural coordinates of at least a portion of a polypeptide of the invention and having a root mean square deviation from the structure coordinates of the polypeptide of the invention of less than 1.50, 1.40, 1.25, 1.0, 0.75, 0.5 or 0.35 Å. In certain embodiments, the portion of a polypeptide of the invention is 25%, 33%, 50%, 66%, 75%,

85%, 90%, or 95% or more of the amino acid residues contained in the polypeptide.

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In another aspect, the present invention provides a molecule or complex including a druggable region of a polypeptide of the invention, the druggable region being defined by a set of points having a root mean square deviation of less than about 1.75 Å from the structural coordinates for points representing (a) the backbone atoms of the amino acids contained in a druggable region of a polypeptide of the invention, (b) the side chain atoms (and optionally the  $C\alpha$  atoms) of the amino acids contained in such druggable region, or (c) all the atoms of the amino acids contained in such druggable region. In certain embodiments, only a portion of the amino acids of a druggable region can be included in the set of points, such as 25%, 33%, 50%, 66%, 75%, 85%, 90% or 95% or more of the amino acid residues contained in the druggable region. In certain embodiments, the root mean square deviation can be less than 1.50, 1.40, 1.25, 1.0, 0.75, 0.5, or 0.35 Å. In still other embodiments, instead of a druggable region, a stable domain, fragment, or structural motif is used in place of a druggable region.

### VI.A.4. Machine Displays and Machine Readable Storage Media

The invention provides a machine-readable storage medium including a data storage material encoded with machine readable data which, when using a machine programmed with instructions for using said data, displays a graphical three-dimensional representation of any of the molecules or complexes, or portions thereof, of this invention. In another embodiment, the graphical three-dimensional representation of such molecule, complex, or portion thereof includes the root mean square deviation of certain atoms of such molecule by a specified amount, such as the backbone atoms by less than 1.5 Å. In another embodiment, a structural equivalent of such molecule, complex, or portion thereof, can be displayed. In another embodiment, the portion can include a druggable region of the polypeptide of the invention.

According to one embodiment, the invention provides a computer for determining at least a portion of the structure coordinates corresponding to X-

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ray diffraction data obtained from a molecule or complex, wherein said computer includes: (a) a machine-readable data storage medium comprising a data storage material encoded with machine-readable data, wherein said data comprises at least a portion of the structural coordinates of a polypeptide of the invention; (b) a machine-readable data storage medium comprising a data storage material encoded with machine-readable data, wherein said data comprises X-ray diffraction data from said molecule or complex; (c) a working memory for storing instructions for processing said machine-readable data of (a) and (b); (d) a central-processing unit coupled to said working memory and to said machine-readable data storage medium of (a) and (b) for performing a Fourier transform of the machine readable data of (a) and for processing said machine readable data of (b) into structure coordinates; and (e) a display coupled to said central-processing unit for displaying said structure coordinates of said molecule or complex. In certain embodiments, the structural coordinates displayed are structurally equivalent to the structural coordinates of a polypeptide of the invention.

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In an alternative embodiment, the machine-readable data storage medium includes a data storage material encoded with a first set of machine readable data which includes the Fourier transform of the structure coordinates of a polypeptide of the invention or a portion thereof, and which, when using a machine programmed with instructions for using said data, can be combined with a second set of machine readable data including the X-ray diffraction pattern of a molecule or complex to determine at least a portion of the structure coordinates corresponding to the second set of machine readable data.

For example, a system for reading a data storage medium can include a computer including a central processing unit (CPU), a working memory which can be, *i.e.*, random access memory (RAM) or "core" memory, mass storage memory (such as one or more disk drives or CD-ROM drives), one or more display devices (*i.e.*, cathode-ray tube ("CRT") displays, light emitting diode (LED) displays, liquid crystal displays (LCDs), electroluminescent displays, vacuum fluorescent displays, field emission displays (FEDs), plasma

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displays, projection panels, etc.), one or more user input devices (*i.e.*, keyboards, microphones, mice, touch screens, etc.), one or more input lines, and one or more output lines, all of which are interconnected by a conventional bidirectional system bus. The system can be a stand-alone computer, or can be networked (*i.e.*, through local area networks, wide area networks, intranets, extranets, or the internet) to other systems (*i.e.*, computers, hosts, servers, etc.). The system can also include additional computer controlled devices such as consumer electronics and appliances.

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Input hardware can be coupled to the computer by input lines and can be implemented in a variety of ways. Machine-readable data of this invention can be inputted via the use of a modem or modems connected by a telephone line or dedicated data line. Alternatively or additionally, the input hardware can include CD-ROM drives or disk drives. In conjunction with a display terminal, a keyboard can also be used as an input device.

Output hardware can be coupled to the computer by output lines and can similarly be implemented by conventional devices. By way of example, the output hardware can include a display device for displaying a graphical representation of an active site of this invention using a program such as QUANTA as described herein. Output hardware might also include a printer, so that hard copy output can be produced, or a disk drive, to store system output for later use.

In operation, a CPU coordinates the use of the various input and output devices, coordinates data accesses from mass storage devices, accesses to and from working memory, and determines the sequence of data processing steps. A number of programs can be used to process the machine-readable data of this invention. Such programs are discussed in reference to the computational methods of drug discovery as described herein. References to components of the hardware system are included as appropriate throughout the following description of the data storage medium.

Machine-readable storage devices useful in the present invention include, but are not limited to, magnetic devices, electrical devices, optical devices, and combinations thereof. Examples of such data storage devices

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include, but are not limited to, hard disk devices, CD devices, digital video disk devices, floppy disk devices, removable hard disk devices, magneto-optic disk devices, magnetic tape devices, flash memory devices, bubble memory devices, holographic storage devices, and any other mass storage peripheral device. It should be understood that these storage devices include necessary hardware (i.e., drives, controllers, power supplies, etc.) as well as any necessary media (i.e., disks, flash cards, etc.) to enable the storage of data.

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In one embodiment, the present invention contemplates a computer readable storage medium comprising structural data, wherein the data include the identity and three-dimensional coordinates of a polypeptide of the invention or portion thereof. In another aspect, the present invention contemplates a database comprising the identity and three-dimensional coordinates of a polypeptide of the invention or a portion thereof. Alternatively, the present invention contemplates a database comprising a portion or all of the atomic coordinates of a polypeptide of the invention or portion thereof.

## VI.A.5. Structurally Similar Molecules and Complexes

Structural coordinates for a polypeptide of the invention can be used to aid in obtaining structural information about another molecule or complex. This method of the invention allows determination of at least a portion of the three-dimensional structure of molecules or molecular complexes that contain one or more structural features that are similar to structural features of a polypeptide of the invention. Similar structural features can include, for example, regions of amino acid identity, conserved active site or binding site motifs, and similarly arranged secondary structural elements (*i.e.*,  $\alpha$  helices and  $\exists$  sheets). Many of the methods described above for determining the structure of a polypeptide of the invention can be used for this purpose as well.

For the present invention, a "structural homolog" is a polypeptide that contains one or more amino acid substitutions, deletions, additions, or rearrangements with respect to the amino acid sequence of SEQ ID NOs: 2

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or 4 or other polypeptide of the invention, but that, when folded into its native conformation, exhibits or is reasonably expected to exhibit at least a portion of the tertiary (three-dimensional) structure of the polypeptide encoded by SEQ ID NOs: 2 or 4 or such other polypeptide of the invention. For example, structurally homologous molecules can contain deletions or additions of one or more contiguous or noncontiguous amino acids, such as a loop or a domain. Structurally homologous molecules also include modified polypeptide molecules that have been chemically or enzymatically derivatized at one or more constituent amino acids, including side chain modifications, backbone modifications, and N- and C-terminal modifications including acetylation, hydroxylation, methylation, amidation, and the attachment of carbohydrate or lipid moieties, cofactors, and the like.

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By using molecular replacement, all or part of the structure coordinates of a polypeptide of the invention can be used to determine the structure of a crystallized molecule or complex whose structure is unknown more quickly and efficiently than attempting to determine such information *ab initio*. For example, in one embodiment this invention provides a method of utilizing molecular replacement to obtain structural information about a molecule or complex whose structure is unknown including: (a) crystallizing the molecule or complex of unknown structure; (b) generating an X-ray diffraction pattern from said crystallized molecule or complex; and (c) applying at least a portion of the structure coordinates for a polypeptide of the invention to the X-ray diffraction pattern to generate a three-dimensional electron density map of the molecule or complex whose structure is unknown.

In another aspect, the present invention provides a method for generating a preliminary model of a molecule or complex whose structure coordinates are unknown, by orienting and positioning the relevant portion of a polypeptide of the invention within the unit cell of the crystal of the unknown molecule or complex so as best to account for the observed X-ray diffraction pattern of the crystal of the molecule or complex whose structure is unknown.

Structural information about a portion of any crystallized molecule or complex that is sufficiently structurally similar to a portion of a polypeptide of

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the invention can be resolved by this method. In addition to a molecule that shares one or more structural features with a polypeptide of the invention, a molecule that has similar bioactivity, such as the same catalytic activity, substrate specificity or ligand-binding activity as a polypeptide of the invention, can also be sufficiently structurally similar to a polypeptide of the invention to permit use of the structure coordinates for a polypeptide of the invention to solve its crystal structure.

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In another aspect, the method of molecular replacement is utilized to obtain structural information about a complex containing a polypeptide of the invention, such as a complex between a modulator and a polypeptide of the invention (or a domain, fragment, ortholog, homolog etc. thereof). In certain instances, the complex includes a polypeptide of the invention (or a domain, fragment, ortholog, homolog etc. thereof) co-complexed with a modulator. For example, in one embodiment, the present invention contemplates a method for making a crystallized complex comprising a polypeptide of the invention, or a fragment thereof, and a compound having a molecular weight of less than 5 kDa, the method comprising: (a) crystallizing a polypeptide of the invention such that the crystals will diffract X-rays to a resolution of 2.5 Å or better; and (b) soaking the crystal in a solution comprising the compound having a molecular weight of less than 5 kDa, thereby producing a crystallized complex comprising the polypeptide and the compound.

Using homology modeling, a computer model of a structural homolog or other polypeptide can be built or refined without crystallizing the molecule. For example, in another aspect, the present invention provides a computer-assisted method for homology modeling a structural homolog of a polypeptide of the invention including: aligning the amino acid sequence of a known or suspected structural homolog with the amino acid sequence of a polypeptide of the invention and incorporating the sequence of the homolog into a model of a polypeptide of the invention derived from atomic structure coordinates to yield a preliminary model of the homolog; subjecting the preliminary model to energy minimization to yield an energy minimized model; remodeling regions

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of the energy minimized model where stereochemistry restraints are violated to yield a final model of the homolog.

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In another embodiment, the present invention contemplates a method for determining the crystal structure of a homolog of a polypeptide having SEQ ID NO: 2 or SEQ ID NO: 4, or equivalent thereof, the method comprising: (a) providing the three dimensional structure of a crystallized polypeptide having SEQ ID NO: 2 or SEQ ID NO: 4, or a fragment thereof; (b) obtaining crystals of a homologous polypeptide comprising an amino acid sequence that is at least 80% identical to the amino acid sequence set forth in SEQ ID NO: 2 or SEQ ID NO: 4 such that the three dimensional structure of the crystallized homologous polypeptide can be determined to a resolution of 2.5 Å or better; and (c) determining the three dimensional structure of the crystallized homologous polypeptide by X-ray crystallography based on the atomic coordinates of the three dimensional structure provided in step (a). In certain instances of the foregoing method, the atomic coordinates for the homologous polypeptide have a root mean square deviation from the backbone atoms of the polypeptide having SEQ ID NO: 2 or SEQ ID NO: 4, or a fragment thereof, of not more than 1.5 Å for all backbone atoms shared in common with the homologous polypeptide and the polypeptide having SEQ ID NO: 2 or SEQ ID NO: 4, or a fragment thereof.

In another aspect, the present invention provides a method for building a model for the activated conformation of CAR, using the repressed structure of Table 2 as a template. In one embodiment, the method comprises: (a) taking the coordinates for residues 107 to 332 directly from Table 2, effectively assuming that the conformation of this portion of CAR is similar or identical in the activated and repressed states; (b) rotating and translating an X-ray structure of VDR, the Vitamin-D receptor, so as to superimpose its core backbone atoms onto corresponding atoms from CAR; (c) combining the superimposed VDR AF2 helix, residues 416-423, with residues 107-332 from the initial CAR model of step (a), to serve as the starting model for residues 107-332 and 341-348 of the CAR protein in the activated conformation; (d) computationally mutating Val418, Leu419, Val421, Phe422 and Gly423 in the

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transplanted VDR AF2 helix to the corresponding amino acid types in the CAR AF2 helix, which are Leu343, Gln344, Ile346, Cys347 and Ser348, respectively; and (e) adjusting the conformations of the mutated amino acid side-chains in the AF2 helix of the CAR model, residues 343, 344, and 346-348, to avoid overlaps by using either manual manipulation within molecular graphics programs or conformational search and energy minimization. In one embodiment, the method further comprises modeling the CAR AF2 linker region, residues 333-340, by using a computational loop modeling technique, recognizing that the calculated linker conformation would probably deviate considerably from the actual linker conformation.

#### VII. Formation of CAR Ligand-Binding Domain-Ligand Crystals

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The present invention provides crystals of CAR LBD in complex with the ligand. The crystals were obtained using the methodology disclosed in the Examples. The CAR LBD-ligand crystals, which can be native or derivative crystals, have orthorhombic unit cells (an orthorhombic unit cell is a unit cell wherein a  $\neq$  b  $\neq$  c, and wherein  $\alpha$  =  $\beta$  =  $\gamma$  = 90°) and space group symmetry P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub>. There are four CAR LBD molecules in the asymmetric unit. In this CAR crystalline form, the unit cell has dimensions of a = 83.0 Å, b = 116.8 Å, c = 131.9 Å, and  $\alpha$  =  $\beta$  =  $\gamma$  = 90°. This crystal form can be formed in a crystallization reservoir comprising 1  $\mu$ l of the protein-ligand solutions disclosed herein, and 1  $\mu$ l of well buffer (e.g. 100-400 mM sodium potassium tartrate, pH 7.1-7.4).

The native and derivative co-crystals comprising a CAR LBD and a ligand disclosed in the present invention can be obtained by a variety of techniques, including batch, liquid bridge, dialysis, vapor diffusion and hanging drop methods (see e.g., McPherson, 1982; McPherson, 1990; Weber, 1991). In one embodiment, the vapor diffusion and hanging drop methods are used for the crystallization of CAR polypeptides and fragments thereof.

Native crystals of the present invention can be grown by dissolving a substantially pure CAR polypeptide or a fragment thereof, and optionally a

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ligand, in an aqueous buffer containing a precipitant at a concentration just below that necessary to precipitate the protein. Water is removed by controlled evaporation to produce precipitating conditions, which are maintained until crystal growth ceases.

In one embodiment of the invention, native crystals are grown by vapor diffusion (See e.g., McPherson, 1982; McPherson, 1990). In this method, the polypeptide/precipitant solution is allowed to equilibrate in a closed container with a larger aqueous reservoir having a precipitant concentration optimal for producing crystals. Generally, less than about 25 µL of CAR polypeptide solution is mixed with an equal volume of reservoir solution, giving a precipitant concentration about half that required for crystallization. This solution is suspended as a droplet underneath a coverslip, which is sealed onto the top of the reservoir. The sealed container is allowed to stand until crystals grow. Crystals generally form within two to six weeks, and are suitable for data collection within approximately seven to ten weeks. Of course, those of skill in the art will recognize that the above-described crystallization procedures and conditions can be varied.

#### VIII. Solving a Crystal Structure of the Present Invention

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Crystal structures of the present invention can be solved using a variety of techniques including, but not limited to isomorphous replacement, anomalous scattering, or molecular replacement methods. Computer software packages can also be used to solve a crystal structure of the present invention. Applicable software packages include, but are not limited to X-PLOR™ program (Brünger, 1992; available from Accelrys Inc, San Diego, California, United States of America), Xtal View (McRee, 1992; available from the San Diego Supercomputer Center, San Diego, California, United States of America); SHELXS 97 (Sheldrick, 1990; available from the Institute of Inorganic Chemistry, Georg-August-Universität, Göttingen, Germany); HEAVY (Terwilliger, Los Alamos National Laboratory) and SHAKE-AND-BAKE (Hauptman, 1997; Weeks *et al.*, 1993; available from the Hauptman-

Woodward Medical Research Institute, Buffalo, New York, United States of America). See also, Ducruix & Geige, 1992, and references cited therein.

## IX. The Overall Structure of CARα in Complex With a Ligand

The structure of the LBD of CAR bound with Compound 1 has been determined to 2.15Å. The statistics of the data and the refined structure are summarized in Table 1.

<u>Table 1</u>
<u>Statistics of Crystallographic Data and Structure</u>

Crystals	CAR/ with Compound 1
Space group	P2 <sub>1</sub> 2 <sub>1</sub> 2 <sub>1</sub>
Resolution (Å)	40.0- 2.15
Unique reflections	69,338
Completeness (%)	99.6
l/σ(last shell)	21.7 (3.1)
R <sub>sym</sub> <sup>a</sup> (%)	9.1
Refinement statistics	
R factor <sup>b</sup> (%)	21.5
R free (%)	25.1
R.M.S.D.	
bond lengths (Å)	0.007
R.M.S.D.	
bond angles(degrees)	1.308
Total non-hydrogen atoms	
	8601

R.M.S.D. is the root mean square deviation from ideal geometry.

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 $<sup>{}^{</sup>a}R_{sym} = \sum |lavg - li| / \sum |li|$ 

 $<sup>^{</sup>b}R_{factor} = \sum |F_{P} - F_{Pcalc}| / \sum F_{p}$ , where  $F_{p}$  and  $F_{pcalc}$  are observed and calculated structure factors,  $R_{free}$  is calculated from a randomly chosen 10% of reflections that were never used in refinement and  $R_{factor}$  is calculated for the remaining 90% of reflections.

In its complex with Compound 1, an inverse agonist, the CAR LBD has a structure with approximately 11 alpha helices and a beta-sheet with 3 strands, as shown in Figure 1. The CAR LBD amino acid sequence is more similar to PXR and VDR than to any other NR LBD sequence, with 50% identity to PXR and 40% identity to VDR in a core region corresponding to VDR residues 126-142, 227-289, 293-300, 302-404 and 416-421. Slightly lower percent identities are obtained by considering the entire LBD sequences; however, these percent identities are complicated by the presence of additional amino acids inserted between Helix-1 and Helix-3 in PXR.

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Figure 2 gives an alignment of the human, mouse, and rat CAR sequences with the human PXR and CAR sequences, with annotation and shading to indicate structural features identified from the X-ray structures. The AF2 helix that is normally present in NR LBDs was absent in this structure, but another helix, designated here as "helix-X", was present. Helix-X includes Leu336, Ser337, Ala338, and Met339, which lie between helix-10 and the residues that normally form the AF2 helix. The hydrogen bonding pattern in helix-X is closer to that of a 3-10 helix rather than an ideal alpha helix. The absence of the AF2 helix was initially very surprising, since the amino acid sequence at the C-terminal end of CAR is very similar to the corresponding segments in VDR and PXR (Figure 2), where the AF2 helix has been seen in all available X-ray structures. Normally, activation of gene transcription depends on the binding of a coactivator, such as CREB binding protein (CBP) or steroid receptor coactivator-1 (SRC-1), and this in turn normally requires the presence of the AF2 helix in its active position. Thus, one would expect the AF2 helix to be present and in the active position in the unliganded, constitutively active form of CAR.

An inverse agonist such as Compound 1 or an antagonist could reduce gene transcription by shifting the AF2 helix into an alternative position, as has been observed with estrogen receptor (ER) bound to antagonists such as tamoxifen and raloxifene (Shiau *et al.*, 1998). Alternatively, an inverse agonist

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or antagonist could act by unwinding the AF2 helix without necessarily moving it from its active position. Further analysis of the CAR X-ray structure suggests that helix-X interferes with the formation of the AF2 helix. Also, side-chains from Met339 and Met340, in and adjacent to helix-X, make extensive interactions with Compound 1. This suggests that Compound 1 induces the formation of helix-X, which in turn unwinds the AF2 helix, thereby preventing coactivator binding and shutting down gene transcription.

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More generally, the analysis of the X-ray structure suggests that CAR exists in equilibrium with at least two major conformations. One conformation is an "activated conformation", not yet observed by X-ray crystallography, where the AF2 helix is properly formed and resides in its active position. The second major conformation is an inactivated conformation, exemplified by the complex of CAR with Compound 1, where helix-X is present and the AF2 helix is absent. While the inventors do not wish to be bound by any particular hypothesized mechanism of action, it appears that, in the absence of ligand, CAR exists predominantly in the activated conformation. Agonist and "superagonist" compounds would tend to shift the equilibrium even farther towards this activated form, effectively increasing the fraction of the CAR receptor in the activated state to a level higher than that observed in the absence of ligand. Inverse agonists, such as Compound 1, would act by shifting the equilibrium towards the inactivated conformation, effectively decreasing the fraction of the CAR receptor in the activated state.

The structure of CAR revealed a number of other major structural differences when compared with the structures of PXR and VDR. The CAR X-ray structure allowed an accurate alignment of helix-1, confirming that PXR and VDR have 45 and 51 additional residues, respectively, in the region between helix-1 and helix-3. The conformation of this insert is unknown in VDR, as the available X-ray structures were determined with a construct where this insert was deleted. The full insert was present in the construct used for the PXR X-ray structure, and most of the insert was visible in the electron density. Surprisingly, in PXR, a segment from this insert acts to displace helix-6 from its usual position where it covers the ligand-binding

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pocket. This segment adopts an extended conformation that occupies less volume than helix-6, effectively opening up additional volume for the ligand in the PXR ligand-binding pocket. While the inventors do not wish to be bound by any particular hypothesized mechanism of action, based on the PXR X-ray structure and the similarity of the CAR amino acid sequence to PXR, one might expect that helix-6 would be absent or displaced away from the ligandbinding pocket, and that the ligand-binding pocket would be similarly voluminous. However, the X-ray structure of CAR reveals that helix-6 is present in CAR, and located in a position similar to that in VDR where it serves as one wall for the ligand-binding pocket. This reduces the volume available to the ligand in the ligand-binding pocket, and changes the shape of the pocket substantially. The pocket volume was calculated with the GRASP program using the atomic radii of Bondi, 1964, using a procedure where the MVP program is used to close channels to the external solvent. With this procedure, the CAR pocket has a volume of 824 Å<sup>3</sup>, similar to that of VDR, which has a volume of 871 Å<sup>3</sup> when bound to Vitamin D, but much smaller than PXR, which has a volume of 1150-1544 Å<sup>3</sup>, depending on the ligand complexed to the protein.

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The structure of the LBD of CAR comprises 11 main alpha helices, a beta sheet with 4 strands, and additional irregular structure and shorter helices. The key features are shown in Figure 1. Helices 3, 5, 6, 7, and 10 and beta strands 2, 3, and 4 enclose the ligand-binding pocket, like a three-layer sandwich (Figure 6). Helix 6, which is absent or displaced in PXR, is intact in CAR, and located in a position similar to that in VDR where it serves as part of the wall of the ligand-binding site. The structure-based sequence alignment of Figure 2 shows the secondary structures of CAR, PXR, and VDR. The presence of helix 6 in CAR reduces the size of the ligand-binding site. The limited binding pocket gives more selectivity in ligand-binding in CAR than in PXR. Binding of the antagonist in CAR causes the AF2 helix to unwind. Instead, a short sequence of amino acids located between helix 10 and the AF2 helix (Leu336, Ser337, Ala338, Met339) form a short 3-10 helix. The side chains of Leu336 and Met339, from the 3-10 helix, and Met340 form

a wall that nicely fits the side of the phenyl ring of the ligand (Figure 1 & 3). This 3-10 helix is referred to as helix X. Steric hindrance from helix X appears to contribute to the unwinding of AF2 helix

The ligand-binding site can be divided into two chambers (Figure 5). One chamber contains the phenylethyl and benzimidazole-6-carboxamide fragments of the ligand. It is completely shielded from solvent. The other chamber contains the benzhydryl fragment of the ligand. This chamber is exposed to the solvent. The amino linker of the ligand is near the interface of the two chambers.

Figure 3 and 4 shows that the ligand fits nicely into the hydrophobic pocket of the LBD site formed mostly by aromatic or hydrophobic residues. They are Phe132, Phe161, Ile164, Asn165, Thr166, Met168, Val169, Ala198, Val199, Cys202, His203, Leu206, Phe217, Tyr224, Thr225, Ile226, Glu227, Asp228, Gly229, Ala230, Phe234, Phe238, Leu239, Leu242, Phe243, His246, Tyr326, Ile330, Leu336, Ser337, Met339, and Met340.

As shown in Figure 3 and 4, there are four hydrogen bonds between the ligand and LBD. The benzimidazol-6-carboxamide forms hydrogen bonds with the carbonyl oxygen of Thr225 and Gly229 amide, respectively. The unsubstituted nitrogen on the benzimidazole forms a hydrogen bond with the hydroxyl group of Tyr326. The amino group linked to the benzhydryl forms a hydrogen bond with the carboxyl oxygen of Asn165. The later two hydrogen bonds are located near the intersection of the two chambers.

### X. Rational Drug Design

#### X.A. Generally

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Modulators to polypeptides of the invention and other structurally related molecules, and complexes containing the same, can be identified and developed as set forth below and otherwise using techniques and methods known to those of skill in the art.

30 The present invention contemplates making any molecule that is shown to modulate the activity of a polypeptide of the invention.

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In another embodiment, inhibitors, modulators of the subject polypeptides, or biological complexes containing them, can be used in the manufacture of a medicament for any number of uses, including, for example, treating any disease or other treatable condition of a patient (including humans and animals), and particularly a disease caused by aberrant CAR regulation or activity.

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A number of techniques can be used to screen, identify, select, and design chemical entities capable of associating with polypeptides of the invention, structurally homologous molecules, and other molecules. Knowledge of the structure for a polypeptide of the invention, determined in accordance with the methods described herein, permits the design and/or identification of molecules and/or other modulators which have a shape complementary to the conformation of a polypeptide of the invention, or more particularly, a druggable region thereof. It is understood that such techniques and methods can use, in addition to the exact structural coordinates and other information for a polypeptide of the invention, structural equivalents thereof described above (including, for example, those structural coordinates that are derived from the structural coordinates of amino acids contained in a druggable region as described above).

The term "chemical entity", as used herein, refers to chemical compounds, complexes of two or more chemical compounds, and fragments of such compounds or complexes. In certain instances, it is desirable to use chemical entities exhibiting a wide range of structural and functional diversity, such as compounds exhibiting different shapes (*i.e.*, flat aromatic rings(s), puckered aliphatic rings(s), straight and branched chain aliphatics with single, double, or triple bonds) and diverse functional groups (*i.e.*, carboxylic acids, esters, ethers, amines, aldehydes, ketones, and various heterocyclic rings).

In one aspect, the method of drug design generally includes computationally evaluating the potential of a selected chemical entity to associate with any of the molecules or complexes of the present invention (or portions thereof). For example, this method can include the steps of (a) employing computational means to perform a fitting operation between the

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selected chemical entity and a druggable region of the molecule or complex; and (b) analyzing the results of said fitting operation to quantify the association between the chemical entity and the druggable region.

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A chemical entity can be examined either through visual inspection or through the use of computer modeling using a docking program such as GRAM, DOCK, or AUTODOCK (Dunbrack et al., 1997). This procedure can include computer fitting of chemical entities to a target to ascertain how well the shape and the chemical structure of each chemical entity will complement or interfere with the structure of the subject polypeptide (Bugg et al., 1993; West et al., 1995). Computer programs can also be employed to estimate the attraction, repulsion, and steric hindrance of the chemical entity to a druggable region, for example. Generally, the tighter the fit (i.e., the lower the steric hindrance, and/or the greater the attractive force) the more potent the chemical entity will be because these properties are consistent with a tighter binding constant. Furthermore, the more specificity in the design of a chemical entity the more likely that the chemical entity will not interfere with related proteins, which can minimize potential side-effects due to unwanted interactions.

A variety of computational methods for molecular design, in which the steric and electronic properties of druggable regions are used to guide the design of chemical entities, are known. See e.g., Cohen et al., 1990; Kuntz et al., 1982; DesJarlais, 1988; Bartlett et al., 1989; Goodford et al., 1985; DesJarlais et al., 1986. Directed methods generally fall into two categories: (1) design by analogy in which 3-D structures of known chemical entities (such as from a crystallographic database) are docked to the druggable region and scored for goodness-of-fit; and (2) de novo design, in which the chemical entity is constructed piece-wise in the druggable region. The chemical entity can be screened as part of a library or a database of molecules. Databases which can be used include ACD (MDL Systems Inc., San Leandro, California, United States of America), NCI (National Cancer Institute, Bethesda, Maryland, United States of America), CCDC (Cambridge Crystallographic Data Center, Cambridge, England, United Kingdom), CAST

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(Chemical Abstract Service), Derwent (Derwent Information Limited, London, England, United Kingdom), Maybridge (Maybridge Chemical Company Ltd., Cornwall, England, United Kingdom), Aldrich (Aldrich Chemical Company, St. Louis, Missouri, United States of America), DOCK (University of California in San Francisco, San Francisco, California, United States of America), and the Directory of Natural Products (Chapman & Hall). Computer programs such as CONCORD (Tripos Inc., St. Louis, Missouri, United States of America) or DB-Converter (Molecular Simulations Limited, Cambridge, England, United Kingdom) can be used to convert a data set represented in two dimensions to one represented in three dimensions.

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Chemical entities can be tested for their capacity to fit spatially with a druggable region or other portion of a target protein. As used herein, the term "fits spatially" means that the three-dimensional structure of the chemical entity is accommodated geometrically by a druggable region. A favorable geometric fit occurs when the surface area of the chemical entity is in close proximity with the surface area of the druggable region without forming unfavorable interactions. A favorable complementary interaction occurs where the chemical entity interacts by hydrophobic, aromatic, ionic, dipolar, or hydrogen donating and accepting forces. Unfavorable interactions can be steric hindrance between atoms in the chemical entity and atoms in the druggable region.

If a model of the present invention is a computer model, the chemical entities can be positioned in a druggable region through computational docking. If, on the other hand, the model of the present invention is a structural model, the chemical entities can be positioned in the druggable region by, for example, manual docking. As used herein the term "docking" refers to a process of placing a chemical entity in close proximity with a druggable region, or a process of finding low energy conformations of a chemical entity/druggable region complex.

In an illustrative embodiment, the design of potential modulator begins from the general perspective of shape complimentary for the druggable region of a polypeptide of the invention, and a search algorithm is employed which is

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capable of scanning a database of small molecules of known three-dimensional structure for chemical entities which fit geometrically with the target druggable region. Most algorithms of this type provide a method for finding a wide assortment of chemical entities that are complementary to the shape of a druggable region of the subject polypeptide. Each of a set of chemical entities from a particular data-base, such as the Cambridge Crystallographic Data Bank (CCDB) (Allen et al., 1973), is individually docked to the druggable region of a polypeptide of the invention in a number of geometrically permissible orientations with use of a docking algorithm. In certain embodiments, a set of computer algorithms called DOCK, can be used to characterize the shape of invaginations and grooves that form the active sites and recognition surfaces of the druggable region (Kuntz et al., 1982). The program can also search a database of small molecules for templates whose shapes are complementary to particular binding sites of a polypeptide of the invention (DesJarlais et al., 1988).

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The orientations are evaluated for goodness-of-fit and the best are kept for further examination using molecular mechanics programs, such as AMBER or CHARMM. Such algorithms have previously proven successful in finding a variety of chemical entities that are complementary in shape to a druggable region.

Goodford et al., 1985 and Boobbyer et al., 1989 have produced a computer program (GRID) that seeks to determine regions of high affinity for different chemical groups (termed probes) of the druggable region. GRID hence provides a tool for suggesting modifications to known chemical entities that might enhance binding. It can be anticipated that some of the sites discerned by GRID as regions of high affinity correspond to "pharmacophoric patterns" determined inferentially from a series of known ligands. As used herein, a "pharmacophoric pattern" is a geometric arrangement of features of chemical entities that is believed to be important for binding. Attempts have been made to use pharmacophoric patterns as a search screen for novel ligands (Jakes et al., 1987; Brint & Willett, 1987; Jakes et al., 1986).

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Yet a further embodiment of the present invention utilizes a computer algorithm such as CLIX which searches such databases as CCDB for chemical entities which can be oriented with the druggable region in a way that is both sterically acceptable and has a high likelihood of achieving favorable chemical interactions between the chemical entity and the surrounding amino acid residues. The method is based on characterizing the region in terms of an ensemble of favorable binding positions for different chemical groups and then searching for orientations of the chemical entities that cause maximum spatial coincidence of individual candidate chemical groups with members of the ensemble. The algorithmic details of CLIX are described in Lawrence et al., 1992.

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In this way, the efficiency with which a chemical entity can bind to or interfere with a druggable region can be tested and optimized by computational evaluation. For example, for a favorable association with a druggable region, a chemical entity must preferably demonstrate a relatively small difference in energy between its bound and fine states (*i.e.*, a small deformation energy of binding). Thus, certain, more desirable chemical entities will be designed with a deformation energy of binding of not greater than about 10 kcal/mole, and more preferably, not greater than 7 kcal/mole. Chemical entities can interact with a druggable region in more than one conformation that is similar in overall binding energy. In those cases, the deformation energy of binding is taken to be the difference between the energy of the free entity and the average energy of the conformations observed when the chemical entity binds to the target.

In this way, the present invention provides computer-assisted methods for identifying or designing a potential modulator of the activity of a polypeptide of the invention including: supplying a computer modeling application with a set of structure coordinates of a molecule or complex, the molecule or complex including at least a portion of a druggable region from a polypeptide of the invention; supplying the computer modeling application with a set of structure coordinates of a chemical entity; and determining whether the chemical entity is expected to bind to the molecule or complex, wherein

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binding to the molecule or complex is indicative of potential modulation of the activity of a polypeptide of the invention.

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In another aspect, the present invention provides a computer-assisted method for identifying or designing a potential modulator to a polypeptide of the invention, supplying a computer modeling application with a set of structure coordinates of a molecule or complex, the molecule or complex including at least a portion of a druggable region of a polypeptide of the invention; supplying the computer modeling application with a set of structure coordinates for a chemical entity; evaluating the potential binding interactions between the chemical entity and active site of the molecule or molecular complex; structurally modifying the chemical entity to yield a set of structure coordinates for a modified chemical entity, and determining whether the modified chemical entity is expected to bind to the molecule or complex, wherein binding to the molecule or complex is indicative of potential modulation of the polypeptide of the invention.

In one embodiment, a potential modulator can be obtained by screening a peptide library (Scott & Smith, 1990; Cwirla et al., 1990; Devlin et al., 1990). A potential modulator selected in this manner could then be systematically modified by computer modeling programs until one or more promising potential drugs are identified. Such analysis has been shown to be effective in the development of HIV protease inhibitors (Lam et al., 1994; Wlodawer et al., 1993; Appelt, 1993; Erickson, 1993). Alternatively a potential modulator can be selected from a library of chemicals such as those that can be licensed from third parties, such as chemical and pharmaceutical companies. A third alternative is to synthesize the potential modulator de novo.

For example, in certain embodiments, the present invention provides a method for making a potential modulator for a polypeptide of the invention, the method including synthesizing a chemical entity or a molecule containing the chemical entity to yield a potential modulator of a polypeptide of the invention, the chemical entity having been identified during a computer-assisted process including supplying a computer modeling application with a set of structure

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coordinates of a molecule or complex, the molecule or complex including at least one druggable region from a polypeptide of the invention; supplying the computer modeling application with a set of structure coordinates of a chemical entity; and determining whether the chemical entity is expected to bind to the molecule or complex at the active site, wherein binding to the molecule or complex is indicative of potential modulation. This method can further include the steps of evaluating the potential binding interactions between the chemical entity and the active site of the molecule or molecular complex and structurally modifying the chemical entity to yield a set of structure coordinates for a modified chemical entity, which steps can be repeated one or more times.

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Once a potential modulator is identified, it can then be tested in any standard assay for the macromolecule depending of course on the macromolecule, including in high throughput assays. Further refinements to the structure of the modulator will generally be necessary and can be made by the successive iterations of any and/or all of the steps provided by the particular screening assay, in particular further structural analysis by *i.e.*, 15N NMR relaxation rate determinations or X-ray crystallography with the modulator bound to the subject polypeptide. These studies can be performed in conjunction with biochemical assays.

Once identified, a potential modulator can be used as a model structure, and analogs to the compound can be obtained. The analogs are then screened for their ability to bind the subject polypeptide. An analog of the potential modulator might be chosen as a modulator when it binds to the subject polypeptide with a higher binding affinity than the predecessor modulator.

In a related approach, iterative drug design is used to identify modulators of a target protein. Iterative drug design is a method for optimizing associations between a protein and a modulator by determining and evaluating the three dimensional structures of successive sets of protein/modulator complexes. In iterative drug design, crystals of a series of protein/modulator complexes are obtained and then the three-dimensional

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structures of each complex is solved. Such an approach provides insight into the association between the proteins and modulators of each complex. For example, this approach can be accomplished by selecting modulators with inhibitory activity, obtaining crystals of this new protein/modulator complex, solving the three dimensional structure of the complex, and comparing the associations between the new protein/modulator complex and previously solved protein/modulator complexes. By observing how changes in the modulator affected the protein/modulator associations, these associations can be optimized.

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In addition to designing and/or identifying a chemical entity to associate with a druggable region, as described above, the same techniques and methods can be used to design and/or identify chemical entities that either associate, or do not associate, with affinity regions, selectivity regions or undesired regions of protein targets. By such methods, selectivity for one or a few targets, or alternatively for multiple targets, from the same species or from multiple species, can be achieved.

For example, a chemical entity can be designed and/or identified for which the binding energy for one druggable region, *i.e.*, an affinity region or selectivity region, is more favorable than that for another region, *i.e.*, an undesired region, by about 20%, 30%, 50% to about 60% or more. It can be the case that the difference is observed between (a) more than two regions, (b) between different regions (selectivity, affinity or undesirable) from the same target, (c) between regions of different targets, (d) between regions of homologs from different species, or (e) between other combinations. Alternatively, the comparison can be made by reference to the  $K_d$ , usually the apparent  $K_d$ , of said chemical entity with the two or more regions in question.

In another aspect, prospective modulators are screened for binding to two nearby druggable regions on a target protein. For example, a modulator that binds a first region of a target polypeptide does not bind a second nearby region. Binding to the second region can be determined by monitoring changes in a different set of amide chemical shifts in either the original screen or a second screen conducted in the presence of a modulator (or potential

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modulator) for the first region. From an analysis of the chemical shift changes, the approximate location of a potential modulator for the second region is identified. Optimization of the second modulator for binding to the region is then carried out by screening structurally related compounds (*i.e.*, analogs as described above).

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When modulators for the first region and the second region are identified, their location and orientation in the ternary complex can be determined experimentally. On the basis of this structural information, a linked compound, i.e., a consolidated modulator, is synthesized in which the modulator for the first region and the modulator for the second region are linked. In certain embodiments, the two modulators are covalently linked to form a consolidated modulator. This consolidated modulator can be tested to determine if it has a higher binding affinity for the target than either of the two individual modulators. A consolidated modulator is selected as a modulator when it has a higher binding affinity for the target than either of the two modulators. Larger consolidated modulators can be constructed in an analogous manner, i.e., linking three modulators which bind to three nearby regions on the target to form a multilinked consolidated modulator that has an even higher affinity for the target than the linked modulator. In this example, it is assumed that is desirable to have the modulator bind to all the druggable regions. However, it can be the case that binding to certain of the druggable regions is not desirable, so that the same techniques can be used to identify modulators and consolidated modulators that show increased specificity based on binding to at least one but not all druggable regions of a target.

The present invention provides a number of methods that use drug design as described above. For example, in one aspect, the present invention contemplates a method for designing a candidate compound for screening for inhibitors of a polypeptide of the invention, the method comprising: (a) determining the three dimensional structure of a crystallized polypeptide of the invention or a fragment thereof; and (b) designing a candidate inhibitor based on the three dimensional structure of the crystallized polypeptide or fragment.

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In another aspect, the present invention provides a method for identifying a potential inhibitor of a polypeptide of the invention, the method comprising: (a) providing the three-dimensional coordinates of a polypeptide of the invention or a fragment thereof; (b) identifying a druggable region of the polypeptide or fragment; and (c) selecting from a database at least one compound that comprises three dimensional coordinates which indicate that the compound can bind the druggable region; (d) wherein the selected compound is a potential inhibitor of a polypeptide of the invention.

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In another aspect, the present invention contemplates a method for identifying a potential modulator of a molecule comprising a druggable region similar to that of SEQ ID NO: 2 or SEQ ID NO: 4, the method comprising: (a) using the atomic coordinates of amino acid residues from SEQ ID NO: 2 or SEQ ID NO: 4, or a fragment thereof, ± a root mean square deviation from the backbone atoms of the amino acids of not more than 1.5 Å, to generate a three-dimensional structure of a molecule comprising a druggable region that is a portion of SEQ ID NO: 2 or SEQ ID NO: 4; (b) employing the three dimensional structure to design or select the potential modulator; (c) synthesizing the modulator; and (d) contacting the modulator with the molecule to determine the ability of the modulator to interact with the molecule.

In another aspect, the present invention contemplates an apparatus for determining whether a compound is a potential inhibitor of a polypeptide having SEQ ID NO: 2 or SEQ ID NO: 4, the apparatus comprising: (a) a memory that comprises: (i) the three dimensional coordinates and identities of the atoms of a polypeptide of the invention or a fragment thereof that form a druggable site; and (ii) executable instructions; and (b) a processor that is capable of executing instructions to: (i) receive three-dimensional structural information for a candidate compound; (ii) determine if the three-dimensional structure of the candidate compound is complementary to the structure of the interior of the druggable site; and (iii) output the results of the determination.

In another aspect, the present invention contemplates a method for designing a potential compound for the prevention or treatment of a disease

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or disorder, the method comprising: (a) providing the three dimensional structure of a crystallized polypeptide of the invention, or a fragment thereof; (b) synthesizing a potential compound for the prevention or treatment of a disease or disorder based on the three dimensional structure of the crystallized polypeptide or fragment; (c) contacting a polypeptide of the present invention or a PDE with the potential compound; and (d) assaying the activity of a polypeptide of the present invention, wherein a change in the activity of the polypeptide indicates that the compound can be useful for prevention or treatment of a disease or disorder.

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In another aspect, the present invention contemplates a method for designing a potential compound for the prevention or treatment of a disease or disorder, the method comprising: (a) providing structural information of a druggable region derived from NMR spectroscopy of a polypeptide of the invention, or a fragment thereof; (b) synthesizing a potential compound for the prevention or treatment of a disease or disorder based on the structural information; (c) contacting a polypeptide of the present invention or a PDE with the potential compound; and (d) assaying the activity of a polypeptide of the present invention, wherein a change in the activity of the polypeptide indicates that the compound can be useful for prevention or treatment of a disease or disorder.

# X.B. Methods of Designing CAR LBD Ligand Compounds

As discussed above, the analysis of the CAR X-ray structure suggests that CAR can adopt at least two major conformations. One major conformation corresponds to the activated state of CAR, where helix-X is absent, and where the AF2 helix is properly formed and resides in its active position. The second major conformation corresponds to the inactivated conformation, exemplified by the complex of CAR with Compound 1, where helix-X is present and where the AF2 helix is absent. In both conformations, the ligand-binding pocket is capped by the C-terminal tail, residues 340-348. These residues adopt different conformations in the activated and inactivated states of CAR, effectively covering the pocket with a cap that can assume at

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least two alternative shapes. Some CAR ligands might bind preferentially to the activated conformation of CAR, whereas some other CAR ligands might bind preferentially to the inactivated conformation of CAR. There might also be some ligands that bind equally well to either conformation of CAR. When a ligand binds preferentially to a particular conformational state, it will lower the energy of that state, thereby shifting the equilibrium towards that state, and increasing the fraction of the CAR receptor that exists in that state. This thermodynamic principle can be used together with the three dimensional structure of CAR to design chemical compounds that bind to specific conformational states of CAR, thereby increasing or decreasing the level of transcription in genes regulated by CAR.

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The present X-ray structure of CAR bound to Compound 1 provides an accurate three-dimensional structure of the ligand-binding pocket in the inactivated conformational state of CAR. Novel ligands can be designed to fit this specific pocket using a variety of computational methods, discussed below. Alternatively, known ligands can be docked into the ligand-binding pocket, using a variety of docking programs and algorithms. These docked structures can be examined graphically to suggest chemical modifications that would improve their fit to the pocket, or their binding to the receptor. Alternatively, known ligands can be complexed with the CAR protein and crystallized using the methods of this invention, allowing the structure of the complex to be determined by X-ray crystallography. The three dimensional structures can be examined graphically to suggest chemical modifications that would improve their fit to the pocket, or their binding to the receptor.

The present X-ray structure of CAR can also be used as a template to build a three-dimensional model of the structure of the activated form of CAR. For example, residues 107 to 332, corresponding to helix-1 through most of helix-10, are taken to have exactly the same coordinates as in the template CAR structure. The AF2 helix, CAR residues 341-348, is then built using the structure of VDR as the template. The VDR template structure is superimposed onto the CAR structure using standard methods as disclosed herein and as would be apparent to one of ordinary skill in the art after a

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review of the present disclosure. The AF2 helix from VDR, residues 416-423, is then removed from the VDR template and transplanted into the model for CAR, without any adjustment of its coordinates. Five of the residues in the VDR AF2 helix have amino acid types different from the corresponding residues in the CAR AF2 helix. These residues are VDR Val418, Leu419, Val421, Phe422, and Gly423, which correspond to CAR Leu343, Gln344, Ile346, Cys347, and Ser348, respectively. These five residues are computationally "mutated" in the model, to obtain the covalent structure corresponding to the desired amino acids in CAR. The C-terminal Ser348 is further modified to obtain a free carboxylate as normally occurs at the C-terminal end of a protein chain.

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These computational mutations can be carried out using amino acid replacement and builder functionality in molecular graphics programs such as Insight-II, available from Accelrys, or using non-graphical molecular mechanics software such as MVP. The side-chain conformations are then adjusted using computer graphics, such as Insight-II, or other energy-based procedures, such as in MVP, to obtain a reasonable overall fit. It is more difficult to obtain a reasonable conformation for the eight residues in the AF2 linker, CAR residues 333-340. The VDR linker, residues 407-415, cannot be used as the template for the CAR linker because it has nine residues, and because its N-terminal end-point is different from that required in CAR. Likewise, the PXR linker, residues 418-422, is too short to serve as a template for the CAR linker. For structure-based drug design, a conservative approach is to omit the linker residues rather than to model the linker incorrectly. Consequently, in one embodiment the linker, residues 333-340, is omitted from the activated CAR model. This model for the activated state of CAR then provides a binding site for the ligand design processes described elsewhere herein. Specifically, various computer software programs can be used to design novel ligands that would fit the specific pocket in the model for the activated form of CAR. Docking calculations can be used to predict how known CAR activators will bind to the activated form of CAR or to identify other available compounds that might bind. These predicted complex

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structures can then be examined by computer graphics to suggest specific chemical modifications that would enhance the binding to the activated state of CAR.

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To be useful as a therapeutic agent, a chemical compound that acts through CAR must induce the appropriate level of CAR activity in relevant In principle, this can be achieved by adjusting the CAR conformational equilibrium so that appropriate fractions of the CAR protein exist in the activated and inactivated states. This in turn can be achieved with ligands that bind almost exclusively to one or the other of the two major conformational states. The design of ligands that are selective for a specific conformational state is facilitated by consideration of how these ligands might bind to each of the two conformational states. Binding modes can be obtained using docking calculations, and then examined graphically to suggest chemical modifications that would make binding to a particular conformational state either more favorable or less favorable. application of these techniques can yield ligands with the desired level of selectivity for the particular conformational state of CAR, thereby achieving the desired level of CAR activity. Ligands that can bind to both conformational states of the CAR protein can also be designed. This is also facilitated by consideration of how the ligands might bind to each of the two conformational states, using the same approach as discussed above, but this time seeking chemical structures and chemical modifications that would permit binding to both conformational states.

The methods of this invention can also be used to suggest possible chemical modifications of a compound that might reduce or minimize its effect on CAR. This approach can be useful in drug discovery projects aiming to find compounds that modulate the activity of some other target molecule, where modulation of CAR activity is an undesirable side effect. This approach is useful in engineering CAR activity out of other, non-drug molecules. Humans and other animals are exposed to a wide range of different chemical compounds, some of which might act on CAR in an undesirable manner. Such a compound could be complexed with CAR and crystallized using the

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methods of the present invention. The structure could then be determined by X-ray crystallography. Alternatively, the structure of the complex could be predicted computationally using molecular docking software. In this case, compounds that tend to activate CAR would be docked into a model or structure of the activated form of CAR, whereas compounds that tend to reduce the activity of CAR would be docked into a model or structure of an inactivated form of CAR, such as its complex with Compound 1 presented here.

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Whether the structure is obtained by X-ray crystallography or computational methods, the structure would be examined by computer graphics to suggest chemical modifications that would minimize the tendency to bind to CAR. For example, substituents could be introduced onto the compound that would project into volume occupied by the CAR protein. Alternatively, a region of the molecule that binds to a lipophilic region of the CAR binding site could be modified to make it more polar, thus reducing its tendency to bind to CAR. Alternatively, a polar group of the compound that makes a hydrogen bonding interaction with CAR could be identified and modified to an alternative group that fails to make the hydrogen bond. Appropriate chemical modifications can be chosen such that the desirable properties and behavior of the compound would be retained.

The design of candidate substances, also referred to as "compounds" or "candidate compounds", that bind to or modulate nuclear receptor (NR) LBD (for example, CAR LBD) -mediated activity according to the present invention generally involves consideration of two factors. First, the compound must be capable of chemically and structurally associating with a NR LBD. Non-covalent molecular interactions important in the association of a NR LBD with its substrate include hydrogen bonding, van der Waals interactions, and hydrophobic interactions. The interaction between an atom of an LBD amino acid and an atom of an LBD ligand can be made by any force or attraction described in nature. Usually the interaction between the atom of the amino acid and the ligand will be the result of a hydrogen bonding interaction, charge interaction, hydrophobic interaction, van der Waals interaction, or dipole

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interaction. In the case of the hydrophobic interaction, it is recognized that this is not a per se interaction between the amino acid and ligand, but rather the usual result, in part, of the repulsion of water or other hydrophilic groups from a hydrophobic surface. Reducing or enhancing the interaction of the LBD and a ligand can be measured by calculating or testing binding energies, either computationally or using thermodynamic or kinetic methods known in the art.

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Second, the compound must be able to assume a conformation that allows it to associate with a NR LBD. Although certain portions of the compound will not directly participate in this association with a NR LBD, those portions can still influence the overall conformation of the molecule. This influence on conformation, in turn, can have a significant impact on potency. Such conformational requirements include the overall three-dimensional structure and orientation of the chemical entity or compound in relation to all or a portion of the binding site, e.g., the ligand-binding pocket or an accessory binding site of a NR LBD, or the spacing between functional groups of a compound comprising several chemical entities that directly interact with a NR LBD.

Chemical modifications can enhance or reduce interactions of an atom of a LBD amino acid and an atom of an LBD ligand. Steric hindrance can be a common approach for changing the interaction of a LBD binding pocket with an activation domain. Chemical modifications are introduced in one embodiment at C-H, C-, and C-OH positions in a ligand, where the carbon is part of the ligand structure that remains the same after modification is complete. In the case of C-H, C could have 1, 2, or 3 hydrogens, but usually only one hydrogen will be replaced. The H or OH can be removed after modification is complete and replaced with a desired chemical moiety.

The potential binding effect of a chemical compound on a NR LBD can be analyzed prior to its actual synthesis and testing by the use of computer modeling techniques that employ the coordinates of a crystalline NR LBD, for example a CAR LBD polypeptide of the present invention. If the theoretical structure of the given compound suggests insufficient interaction and

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association between it and a NR LBD, synthesis and testing of the compound is obviated. However, if computer modeling indicates a strong interaction, the molecule can then be synthesized and tested for its ability to bind and modulate the activity of a NR LBD. In this manner, synthesis of unproductive or inactive compounds can be avoided.

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A binding compound of a NR LBD polypeptide (in one embodiment a CAR LBD) can be computationally evaluated and designed via a series of steps in which chemical entities or fragments are screened and selected for their ability to associate with an individual binding site or other area of a crystalline CAR LBD polypeptide of the present invention and to interact with the amino acids disposed in the binding sites.

Interacting amino acids forming contacts with a ligand and the atoms of the interacting amino acids are usually 2 to 4 angstroms away from the center of the atoms of the ligand. Generally these distances are determined by computer as discussed herein and in McRee, 1993. However distances can be determined manually once the three dimensional model is made. More commonly, the atoms of the ligand and the atoms of interacting amino acids are 3 to 4 angstroms apart. A ligand can also interact with distant amino acids, after chemical modification of the ligand to create a new ligand. Distant amino acids are generally not in contact with the ligand before chemical modification. A chemical modification can change the structure of the ligand to make a new ligand that interacts with a distant amino acid usually at least 4.5 angstroms away from the ligand. Distant amino acids rarely line the surface of the binding cavity for the ligand, as they are too far away from the ligand to be part of a pocket or surface of the binding cavity.

A compound designed or selected as binding to an NR polypeptide (in one embodiment a CAR LBD polypeptide) can be further computationally optimized so that in its bound state it would lack repulsive electrostatic interaction with the target polypeptide. Such non-complementary (e.g., electrostatic) interactions include repulsive charge-charge, dipole-dipole, and charge-dipole interactions. Specifically, the sum of all electrostatic interactions between the ligand and the polypeptide when the ligand is bound

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to an NR LBD make a neutral or favorable contribution to the enthalpy of binding.

One of several methods can be used to screen chemical entities or fragments for their ability to associate with a NR LBD and, more particularly, with the individual binding sites of a NR LBD, such as a ligand-binding pocket or an accessory binding site. This process can begin by visual inspection of, for example, a ligand-binding pocket on a computer screen based on the CAR LBD atomic coordinates disclosed in Tables 2-3. Selected fragments or chemical entities can then be positioned in a variety of orientations, or docked, within an individual binding site of a CAR LBD as defined herein above. Docking can be accomplished using software programs such as those available under the trade names QUANTA<sup>TM</sup> (available from Accelrys Inc, San Diego, California, United States of America) and SYBYL<sup>TM</sup> (available from Tripos, Inc., St. Louis, Missouri, United States of America), followed by energy minimization and molecular dynamics with standard molecular mechanics force fields, such as CHARM (Brooks *et al.*, 1993) and AMBER 5 (Case *et al.*, 1997; Pearlman *et al.*, 1995).

Specialized computer programs can also assist in the process of selecting fragments or chemical entities. These include:

- 1. GRID™ program, version 17 (Goodford, 1985), which is available from Molecular Discovery Ltd. of Oxford, United Kingdom;
- 2. MCSS™ program (Miranker & Karplus, 1991), which is available from Accelrys Inc, San Diego, California, United States of America;
- 3. AUTODOCK™ 3.0 program (Goodsell & Olsen, 1990), which is available from the Scripps Research Institute, La Jolla, California, United States of America;
- 4. DOCK™ 4.0 program (Kuntz et al., 1992), which is available from the University of California, San Francisco, California, United States of America;
- 5. FLEX-X™ program (See Rarey et al., 1996), which is available from Tripos, Inc., St. Louis, Missouri, United States of America;
  - 6. MVP program (Lambert, 1997); and

7. LUDI™ program (Bohm, 1992), which is available from Accelrys Inc, San Diego, California, United States of America.

Once suitable chemical entities or fragments have been selected, they can be assembled into a single compound or ligand. Assembly can proceed by visual inspection of the relationship of the fragments to each other on the three-dimensional image displayed on a computer screen in relation to the structure coordinates of a CAR LBD in complex with a co-regulator, optionally in further complex with a ligand. Manual model building using software such as QUANTA<sup>TM</sup> or SYBYL<sup>TM</sup> typically follows.

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Useful programs to aid one of ordinary skill in the art in connecting the individual chemical entities or fragments include:

- 1. CAVEAT™ program (Bartlett *et al.*, 1989), which is available from the University of California, Berkeley, California, United States of America;
- 2. 3D Database systems, such as MACCS-3D™ system program, which is available from MDL Information Systems, San Leandro, California, United States of America. This area is reviewed in Martin, 1992; and
- 3. HOOK™ program (Eisen *et al.*, 1994), which is available from Accelrys Inc, San Diego, California, United States of America.

Instead of proceeding to build a NR LBD polypeptide ligand (in one embodiment a CAR LBD ligand) in a step-wise fashion one fragment or chemical entity at a time as described above, ligand compounds can be designed as a whole or *de novo* using the structural coordinates of a crystalline CAR LBD polypeptide of the present invention and either an empty binding site or optionally including some portion(s) of a known ligand(s). Applicable methods can employ the following software programs:

- LUDI™ program (Bohm, 1992), which is available from Accelrys Inc,
   San Diego, California, United States of America;
  - 2. LEGEND™ program (Nishibata & Itai, 1991); and
- 3. LEAPFROG™, which is available from Tripos Associates, St. Louis,
   30 Missouri, United States of America.

Other molecular modeling techniques can also be employed in accordance with this invention. See e.g., Cohen et al., 1990; Navia & Murcko,

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1992; and U.S. Patent No. 6,008,033 to <u>Abdel-Meguid *et al.*</u>, all of which are incorporated herein by reference.

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Once a compound has been designed or selected by the above methods, the efficiency with which that compound can bind to a NR LBD can be tested and optimized by computational evaluation. By way of a particular example, a compound that has been designed or selected to function as a CAR LBD ligand can traverse a volume not overlapping that occupied by the binding site when it is bound to its native ligand. Additionally, an effective NR LBD ligand can demonstrate a relatively small difference in energy between its bound and free states (i.e., a small deformation energy of binding). Thus, the most efficient NR LBD ligands can be designed with a deformation energy of binding of in one embodiment not greater than about 10 kcal/mole, and in another embodiment not greater than 7 kcal/mole. It is possible for NR LBD ligands to interact with the polypeptide in more than one conformation that is similar in overall binding energy. In those cases, the deformation energy of binding is taken to be the difference between the energy of the free compound and the thermodynamic average energy of the conformations observed when the ligand binds to the polypeptide.

A compound designed or selected as binding to a NR LBD polypeptide (preferably a CAR polypeptide, more preferably a CAR LBD polypeptide) can be further computationally optimized so that in its bound state it would preferably lack repulsive electrostatic interaction with the target polypeptide. Such non-complementary (e.g., electrostatic) interactions include repulsive charge-charge, dipole-dipole, and charge-dipole interactions. Specifically, the sum of all electrostatic interactions between the ligand and the polypeptide when the ligand is bound to a NR LBD preferably make a neutral or favorable contribution to the enthalpy of binding.

Specific computer software is available in the art to evaluate compound deformation energy and electrostatic interaction. Examples of programs designed for such uses include:

1. GAUSSIAN 98™, which is available from Gaussian, Inc., Pittsburgh, Pennsylvania, United States of America;

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- 2. AMBER™ program, version 6.0, which is available from the University of California, San Francisco, California, United States of America;
- 3. QUANTA™ program, which is available from Accelrys Inc, San Diego, California, United States of America;
- 4. CHARMM® program, which is available from Accelrys Inc, San Diego, California, United States of America; and
- 4. INSIGHT II® program, which is available from Accelrys Inc, San Diego, California, United States of America.

These programs can be implemented using a suitable computer system. Other hardware systems and software packages will be apparent to those skilled in the art after review of the disclosure of the present invention presented herein.

Once a NR LBD modulating compound has been optimally selected or designed, as described above, substitutions can then be made in some of its atoms or side groups in order to improve or modify its binding properties. In some cases, initial substitutions might be conservative, e.g., the replacement group will have approximately the same size, shape, hydrophobicity, and charge as the original group. In other cases, the replacement group will have different properties as desired to make specific interactions with the protein. Such substituted chemical compounds can then be analyzed for efficiency of fit to a NR LBD binding site using the same computer-based approaches described in detail above.

## X.C. Sterically Similar Compounds

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A further aspect of the present invention is that sterically similar compounds can be formulated to mimic the key portions of a CAR LBD structure. Such compounds are functional equivalents. The generation of a structural functional equivalent can be achieved by the techniques of modeling and chemical design known to those of skill in the art and described herein. Modeling and chemical design of CAR and CAR LBD structural equivalents can be based on the structure coordinates of a crystalline CAR

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LBD polypeptide of the present invention. It will be understood that all such sterically similar constructs fall within the scope of the present invention.

## XI. CAR Polypeptides

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The generation of mutant and chimeric CAR polypeptides is also an aspect of the present invention. A chimeric polypeptide can comprise a CAR LBD polypeptide or a portion of a CAR LBD, (e.g. a CAR LBD) which is fused to a candidate polypeptide or a suitable region of the candidate polypeptide. Throughout the present disclosure it is intended that the term "mutant" encompass not only mutants of a CAR LBD polypeptide but chimeric proteins generated using a CAR LBD as well. It is thus intended that the following discussion of mutant CAR LBDs apply mutatis mutandis to chimeric CAR and CAR LBD polypeptides and to structural equivalents thereof.

In accordance with the present invention, a mutation can be directed to a particular site or combination of sites of a wild-type CAR LBD. For example, an accessory binding site or the binding pocket can be chosen for mutagenesis. Similarly, a residue having a location on, at or near the surface of the polypeptide can be replaced, resulting in an altered surface charge of one or more charge units, as compared to the wild-type CAR and CAR LBD. Alternatively, an amino acid residue in a CAR or a CAR LBD can be chosen for replacement based on its hydrophilic or hydrophobic characteristics.

Such mutants can be characterized by any one of several different properties as compared with the wild-type CAR LBD. For example, such mutants can have an altered surface charge of one or more charge units, or can have an increase in overall stability. Other mutants can have altered ligand specificity in comparison with, or a higher specific activity than, a wild type CAR or CAR LBD.

CAR and CAR LBD mutants of the present invention can be generated in a number of ways. For example, the wild-type sequence of a CAR or a CAR LBD can be mutated at those sites identified using this invention as desirable for mutation by employing oligonucleotide-directed mutagenesis or other conventional methods. Alternatively, mutants of a CAR or a CAR LBD

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can be generated by the site-specific replacement of a particular amino acid with an unnaturally occurring amino acid. In addition, CAR or CAR LBD mutants can be generated through replacement of an amino acid residue, for example, a particular cysteine or methionine residue, with selenocysteine or selenomethionine. This can be achieved by growing a host organism capable of expressing either the wild type or mutant polypeptide on a growth medium depleted of either natural cysteine or methionine (or both) but enriched in selenocysteine or selenomethionine (or both).

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Mutations can be introduced into a DNA sequence coding for a CAR or a CAR LBD using synthetic oligonucleotides. These oligonucleotides contain nucleotide sequences flanking the desired mutation sites. Mutations can be generated in the full-length DNA sequence of a CAR or a CAR LBD or in any sequence coding for polypeptide fragments of a CAR or a CAR LBD.

According to the present invention, a mutated CAR or CAR LBD DNA sequence produced by the methods described above, or any alternative methods known in the art, can be expressed using an expression vector. An expression vector, as is well known to those of skill in the art, typically includes elements that permit autonomous replication in a host cell independent of the host genome, and one or more phenotypic markers for selection purposes. Either prior to or after insertion of the DNA sequences surrounding the desired CAR or CAR LBD mutant coding sequence, an expression vector includes control sequences encoding a promoter, operator, ribosome binding site, translation initiation signal, and, optionally, a repressor gene or various activator genes and a signal for termination. Where secretion of the produced mutant is desired, nucleotides encoding a "signal sequence" can be inserted prior to a CAR or a CAR LBD mutant coding sequence. For expression under the direction of the control sequences, a desired DNA sequence is operatively linked to the control sequences; that is, the sequence has an appropriate start signal in front of the DNA sequence encoding the CAR or CAR LBD mutant, and the correct reading frame to permit expression of that sequence under the control of the control sequences and production of the desired product encoded by that CAR or CAR LBD sequence.

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Any of a wide variety of well-known available expression vectors can be used to express a mutated CAR or CAR LBD coding sequences of this invention. These include for example, vectors consisting of segments of chromosomal, non-chromosomal, and synthetic DNA sequences, such as known derivatives of SV40, known bacterial plasmids, e.g., plasmids from E. coli including colE1, pCR1, pBR322, pMB9 and their derivatives, wider host range plasmids, e.g., RP4, phage DNAs, e.g., derivatives of phage λ, e.g., NM 989, and other DNA phages, e.g., M13 and filamentous single stranded DNA phages, yeast plasmids and vectors derived from combinations of plasmids and phage DNAs, such as plasmids which have been modified to employ phage DNA or other expression control sequences. In one embodiment of the present invention, a vector amenable to expression in a pRSETA-based expression system is employed. The pRSETA expression system is available from Invitrogen, Inc., Carlsbad, California, United States of America.

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In addition, any of a wide variety of expression control sequences – *i.e.* sequences that control the expression of a DNA sequence when operatively linked to it – can be used in these vectors to express the mutated DNA sequences according to this invention. Such useful expression control sequences, include, but are not limited to the early and late promoters of SV40 for animal cells; the lac system, the trp system, the TAC or TRC system, the major operator and promoter regions of phage  $\lambda$ , and the control regions of fd coat protein for *E. coli*; the promoter for 3-phosphoglycerate kinase or other glycolytic enzymes, the promoters of acid phosphatase, (for example, Pho5), and the promoters of the yeast  $\alpha$ -mating factors for yeast; as well as other sequences known to control the expression of genes of prokaryotic or eukaryotic cells or their viruses, and various combinations thereof.

A wide variety of hosts can be employed for producing mutated CAR and CAR LBD polypeptides according to this invention. These hosts include, for example, bacteria, such as *E. coli*, *Bacillus*, and *Streptomyces*; fungi, such

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as yeasts; animal cells, such as CHO and COS-1 cells; plant cells; insect cells, such as Sf9 cells; and transgenic host cells.

It should be understood that not all expression vectors and expression systems function in the same way to express mutated DNA sequences of this invention, and to produce modified CAR and CAR LBD polypeptides or CAR or CAR LBD mutants. Neither do all hosts function equally well with the same expression system. One of skill in the art can, however, make a selection among these vectors, expression control sequences and hosts without undue experimentation and without departing from the scope of this invention. For example, an important consideration in selecting a vector will be the ability of the vector to replicate in a given host. The copy number of the vector, the ability to control that copy number, and the expression of any other proteins encoded by the vector, such as antibiotic markers, should also be considered.

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In selecting an expression control sequence, a variety of factors should also be considered. These include, for example, the relative strength of the system, its controllability and its compatibility with the DNA sequence encoding a modified CAR or CAR LBD polypeptide of this invention, with particular regard to the formation of potential secondary and tertiary structures.

Hosts should be selected by consideration of their compatibility with the chosen vector, the toxicity of a modified CAR or CAR LBD to them, their ability to express mature products, their ability to fold proteins correctly, their fermentation requirements, the ease of purification of a modified CAR or CAR LBD and safety. Within these parameters, one of skill in the art can select various vector/expression control system/host combinations that will produce useful amounts of a mutant CAR or CAR LBD. A mutant CAR or CAR LBD produced in these systems can be purified by a variety of conventional steps and strategies, including those used to purify the wild type CAR or CAR LBD.

Once a CAR LBD mutation(s) has been generated in the desired location, such as an active site or dimerization site, the mutants can be tested for any one of several properties of interest. For example, mutants can be screened for an altered charge at physiological pH. This is determined by

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measuring the mutant CAR or CAR LBD isoelectric point (pl) and comparing the observed value with that of the wild-type parent. Isoelectric point can be measured by gel-electrophoresis according to the method of Wellner, 1971. A mutant CAR or CAR LBD polypeptide containing a replacement amino acid located at the surface of the enzyme, as provided by the structural information of this invention, can lead to an altered surface charge and an altered pl.

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## XI.A. Generation of an Engineered CAR LBD or CAR LBD Mutant

In an embodiment of the present invention, a unique CAR or CAR LBD polypeptide is generated. Such a mutant can facilitate purification and the study of the ligand-binding abilities of a CAR polypeptide.

As used in the following discussion, the terms "engineered CAR", "engineered CAR LBD", "CAR mutant", and "CAR LBD mutant" refers to polypeptides having amino acid sequences which contain at least one mutation in the wild-type sequence. The terms also refer to CAR and CAR LBD polypeptides which are capable of exerting a biological effect in that they comprise all or a part of the amino acid sequence of an engineered CAR or CAR LBD polypeptide of the present invention, or cross-react with antibodies raised against an engineered CAR or CAR LBD polypeptide, or retain all or some or an enhanced degree of the biological activity of the engineered CAR or CAR LBD amino acid sequence or protein. Such biological activity can include the binding of small molecules in general, and the binding of Compound 1, in particular.

The terms "engineered CAR LBD" and "CAR LBD mutant" also includes analogs of an engineered CAR LBD or CAR LBD polypeptide. By "analog" is intended that a DNA or polypeptide sequence can contain alterations relative to the sequences disclosed herein, yet retain all or some or an enhanced degree of the biological activity of those sequences. Analogs can be derived from genomic nucleotide sequences or from other organisms, or can be created synthetically. Those of skill in the art will appreciate that other analogs, as yet undisclosed or undiscovered, can be used to design and/or construct CAR LBD or CAR LBD mutant analogs. There is no need for

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a CAR LBD or CAR LBD mutant polypeptide to comprise all or substantially all of the amino acid sequence of SEQ ID NOs: 2 or 4. Shorter or longer sequences can be employed in the invention; shorter sequences are herein referred to as "segments". Thus, the terms "engineered CAR LBD" and "CAR LBD mutant" also includes fusion, chimeric or recombinant CAR LBD or CAR LBD mutant polypeptides and proteins comprising sequences of the present invention. Methods of preparing such proteins are disclosed herein above and are known in the art.

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# XI.A.1. Sequences That Are Substantially Identical to a CAR or CAR LBD Mutant Sequence of the Present Invention

Nucleic acids that are substantially identical to a nucleic acid sequence of a CAR or CAR LBD mutant of the present invention, e.g. allelic variants, genetically altered versions of the gene, etc., bind to a CAR or CAR LBD mutant sequence under stringent hybridization conditions. By using probes, particularly labeled probes of DNA sequences, one can isolate homologous or related genes. The source of homologous genes can be any organism, including, but not limited to primates; rodents, such as rats and mice; canines; felines; bovines; equines; yeast; and nematodes.

Among mammalian species, e.g. human and mouse, homologs can have substantial sequence similarity, i.e. at least 75% sequence identity between nucleotide sequences. Sequence similarity is calculated based on a reference sequence, which can be a subset of a larger sequence, such as a conserved motif, coding region, flanking region, etc. In one embodiment, a reference sequence is at least about 18 nucleotides (nt) long, in another embodiment at least about 30 nt long, and can extend to the complete sequence that is being compared. Algorithms for sequence analysis are known in the art, such as BLAST, described in Altschul et al., 1990.

Percent identity or percent similarity of a DNA or peptide sequence can be determined, for example, by comparing sequence information using the GAP computer program, available from the University of Wisconsin Genetics Computer Group (now part of Accelrys Inc, San Diego, California, United

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States of America). The GAP program utilizes the alignment method of Needleman et al., 1970, as revised by Smith et al., 1981. Briefly, the GAP program defines similarity as the number of aligned symbols (i.e., nucleotides or amino acids) that are similar, divided by the total number of symbols in the shorter of the two sequences. The preferred parameters for the GAP program are the default parameters, which do not impose a penalty for end gaps. See e.g., Schwartz et al., 1979; Gribskov et al., 1986.

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The term "similarity" is contrasted with the term "identity". Similarity is defined as above; "identity", however, refers to a nucleic acid or amino acid sequence having the same amino acid at the same relative position in a given family member of a gene family. Homology and similarity are generally viewed as broader terms than the term identity. Biochemically similar amino acids, for example leucine/isoleucine or glutamate/aspartate, can be present at the same position — these are not identical per se, but are biochemically "similar." As disclosed herein, these are referred to as conservative differences or conservative substitutions. This differs from a conservative mutation at the DNA level, which changes the nucleotide sequence without making a change in the encoded amino acid, e.g. TCC to TCA, both of which encode serine.

As used herein, DNA analog sequences are "substantially identical" to specific DNA sequences disclosed herein if: (a) the DNA analog sequence is derived from coding regions of the nucleic acid sequence shown in SEQ ID NOs: 1 or 3; or (b) the DNA analog sequence is capable of hybridization with DNA sequences of (a) under stringent conditions and which encode a biologically active CAR or CAR LBD gene product; or (c) the DNA sequences are degenerate as a result of alternative genetic code to the DNA analog sequences defined in (a) and/or (b). Substantially identical analog proteins and nucleic acids will have between about 70% and 80%, preferably between about 81% to about 90% or even more preferably between about 91% and 99% sequence identity with the corresponding sequence of the native protein or nucleic acid. Sequences having lesser degrees of identity but comparable biological activity are considered to be equivalents.

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As used herein, "stringent conditions" refers to conditions of high stringency, for example 6X SSC, 0.2% polyvinylpyrrolidone, 0.2% Ficoll, 0.2% bovine serum albumin, 0.1% sodium dodecyl sulfate, 100 μg/ml salmon sperm DNA and 15% formamide at 68°C. For the purposes of specifying additional conditions of high stringency, preferred conditions comprise a salt concentration of about 200 mM and temperature of about 45°C. One example of stringent conditions is hybridization in 4X SSC, at 65°C, followed by a washing in 0.1X SSC at 65°C for one hour. Another exemplary stringent hybridization scheme uses 50% formamide, 4X SSC at 42°C.

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In contrast, nucleic acids having sequence similarity are detected by hybridization under lower stringency conditions. Thus, sequence identity can be determined by hybridization under lower stringency conditions, for example, at 50°C or higher and 0.1X SSC (9 mM NaCl/0.9 mM sodium citrate) and the sequences will remain bound when subjected to washing at 55°C in 1X SSC.

# XI.A.2. Complementarity and Hybridization to an Engineered CAR or CAR LBD Mutant Sequence

As used herein, the term "functionally equivalent codon" is used to refer to codons that encode the same amino acid, such as the ACG and AGU codons for serine. CAR or CAR LBD-encoding nucleic acid sequences comprising SEQ ID NOs: 1 and 3, which have functionally equivalent codons are covered by the present invention. Thus, when referring to the sequence examples presented in SEQ ID NOs: 1 and 3, applicants contemplate substitution of functionally equivalent codons into the sequence example of SEQ ID NOs: 1 and 3. Thus, applicants are in possession of amino acid and nucleic acids sequences which include such substitutions but which are not set forth herein in their entirety for convenience.

It will also be understood by those of skill in the art that amino acid and nucleic acid sequences can include additional residues, such as additional N-or C-terminal amino acids or 5' or 3' nucleic acid sequences, and yet still be essentially as set forth in one of the sequences disclosed herein, so long as

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the sequence retains biological protein activity where polypeptide expression is concerned. The addition of terminal sequences particularly applies to nucleic acid sequences which can, for example, include various non-coding sequences flanking either of the 5' or 3' portions of the coding region or can include various internal sequences, *i.e.*, introns, which are known to occur within genes.

## XI.B. Biological Equivalents

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The present invention envisions and includes biological equivalents of CAR or CAR LBD mutant polypeptide of the present invention. The term "biological equivalent" refers to proteins having amino acid sequences which are substantially identical to the amino acid sequence of a CAR LBD mutant of the present invention and which are capable of exerting a biological effect in that they are capable of binding a small molecule, binding a co-regulator, homo- or heterodimerizing or cross-reacting with anti-CAR or CAR LBD mutant antibodies raised against a mutant CAR or CAR LBD polypeptide of the present invention.

For example, certain amino acids can be substituted for other amino acids in a protein structure without appreciable loss of interactive capacity with, for example, structures in the nucleus of a cell. Since it is the interactive capacity and nature of a protein that defines that protein's biological functional activity, certain amino acid sequence substitutions can be made in a protein sequence (or the nucleic acid sequence encoding it) to obtain a protein with the same, enhanced, or antagonistic properties. Such properties can be achieved by interaction with the normal targets of the protein, but this need not be the case, and the biological activity of the invention is not limited to a particular mechanism of action. It is thus in accordance with the present invention that various changes can be made in the amino acid sequence of a CAR or CAR LBD mutant polypeptide of the present invention or its underlying nucleic acid sequence without appreciable loss of biological utility or activity.

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Biologically equivalent polypeptides, as used herein, are polypeptides in which certain, but not most or all, of the amino acids can be substituted. Thus, when referring to the sequence examples presented in SEQ ID NOs: 2 and 4, applicants envision substitution of codons that encode biologically equivalent amino acids, as described herein, into the sequence example of SEQ ID NOs: 2 and 4, respectively. Thus, applicants are in possession of amino acid and nucleic acids sequences which include such substitutions but which are not set forth herein in their entirety for convenience.

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Alternatively, functionally equivalent proteins or peptides can be created via the application of recombinant DNA technology, in which changes in the protein structure can be engineered, based on considerations of the properties of the amino acids being exchanged, e.g. substitution of Ile for Leu. Changes designed by man can be introduced through the application of site-directed mutagenesis techniques, e.g., to introduce improvements to the antigenicity of the protein or to test a CAR or CAR LBD mutant polypeptide of the present invention in order to modulate co-regulator-binding or other activity, at the molecular level.

Amino acid substitutions, such as those which might be employed in modifying a CAR or CAR LBD mutant polypeptide of the present invention are generally, but not necessarily, based on the relative similarity of the amino acid side-chain substituents, for example, their hydrophobicity, hydrophilicity, charge, size, and the like. An analysis of the size, shape and type of the amino acid side-chain substituents reveals that arginine, lysine and histidine are all positively charged residues; that alanine, glycine and serine are all of similar size; and that phenylalanine, tryptophan and tyrosine all have a generally similar shape. Therefore, based upon these considerations, arginine. lysine and histidine; alanine, glycine and serine; and phenylalanine, tryptophan and tyrosine; are defined herein as biologically functional Those of skill in the art will appreciate other biologically eguivalents. functional equivalent changes. It is implicit in the above discussion, however, that one of skill in the art can appreciate that a radical, rather than a conservative substitution is warranted in a given situation. Non-conservative

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substitutions in mutant CAR or CAR LBD polypeptides of the present invention are also an aspect of the present invention.

In making biologically functional equivalent amino acid substitutions, the hydropathic index of amino acids can be considered. Each amino acid has been assigned a hydropathic index on the basis of their hydrophobicity and charge characteristics, these are: isoleucine (+ 4.5); valine (+ 4.2); leucine (+ 3.8); phenylalanine (+ 2.8); cysteine (+ 2.5); methionine (+ 1.9); alanine (+ 1.8); glycine (-0.4); threonine (-0.7); serine (-0.8); tryptophan (-0.9); tyrosine (-1.3); proline (-1.6); histidine (-3.2); glutamate (-3.5); glutamine (-3.5); aspartate (-3.5); asparagine (-3.5); lysine (-3.9); and arginine (-4.5).

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The importance of the hydropathic amino acid index in conferring interactive biological function on a protein is generally understood in the art (Kyte & Doolittle, 1982, incorporated herein by reference). It is known that certain amino acids can be substituted for other amino acids having a similar hydropathic index or score and still retain a similar biological activity. In making changes based upon the hydropathic index, the substitution of amino acids whose hydropathic indices are within ±2 of the original value is preferred, those within ±1 of the original value are particularly preferred.

It is also understood in the art that the substitution of like amino acids can be made effectively on the basis of hydrophilicity. U.S. Patent No. 4,554,101, incorporated herein by reference, states that the greatest local average hydrophilicity of a protein, as governed by the hydrophilicity of its adjacent amino acids, correlates with its immunogenicity and antigenicity, *i.e.* with a biological property of the protein. It is understood that an amino acid can be substituted for another having a similar hydrophilicity value and still obtain a biologically equivalent protein.

As detailed in U.S. Patent No. 4,554,101 to <u>Hopp</u>, the following hydrophilicity values have been assigned to amino acid residues: arginine (+ 3.0); lysine (+ 3.0); aspartate (+  $3.0\pm1$ ); glutamate (+  $3.0\pm1$ ); serine (+ 0.3); asparagine (+ 0.2); glutamine (+ 0.2); glycine (0); threonine (-0.4); proline (-  $0.5\pm1$ ); alanine (-0.5); histidine (-0.5); cysteine (-1.0); methionine (-1.3); valine

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(-1.5); leucine (-1.8); isoleucine (-1.8); tyrosine (-2.3); phenylalanine (-2.5); tryptophan (-3.4).

In making changes based upon similar hydrophilicity values, the substitution of amino acids whose hydrophilicity values are within ±2 of the original value is preferred, those that are within ±1 of the original value are particularly preferred, and those within ±0.5 of the original value are even more particularly preferred.

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While discussion has focused on functionally equivalent polypeptides arising from amino acid changes, it will be appreciated that these changes can be effected by alteration of the encoding DNA, taking into consideration also that the genetic code is degenerate and that two or more codons can code for the same amino acid.

Thus, it will also be understood that this invention is not limited to the particular amino acid and nucleic acid sequences of SEQ ID NOs: 1-4. Recombinant vectors and isolated DNA segments can therefore variously include a CAR or CAR LBD mutant polypeptide-encoding region itself, include coding regions bearing selected alterations or modifications in the basic coding region, or include larger polypeptides which nevertheless comprise a CAR or CAR LBD mutant polypeptide-encoding regions or can encode biologically functional equivalent proteins or polypeptides which have variant amino acid sequences. Biological activity of a CAR or CAR LBD mutant polypeptide can be determined, for example, by employing binding assays known to those of skill in the art.

The nucleic acid segments of the present invention, regardless of the length of the coding sequence itself, can be combined with other DNA sequences, such as promoters, enhancers, polyadenylation signals, additional restriction enzyme sites, multiple cloning sites, other coding segments, polyhistidine encoding segments and the like, such that their overall length can vary considerably. It is therefore contemplated that a nucleic acid fragment of almost any length can be employed, with the total length preferably being limited by the ease of preparation and use in the intended recombinant DNA protocol. For example, nucleic acid fragments can be

prepared which include a short stretch complementary to a nucleic acid sequence set forth in SEQ ID NOs: 1 and 3, such as about 10 nucleotides, and which are up to 10,000 or 5,000 base pairs in length. DNA segments with total lengths of about 4,000, 3,000, 2,000, 1,000, 500, 200, 100, and about 50 base pairs in length are also useful.

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The DNA segments of the present invention encompass biologically functional equivalents of CAR or CAR LBD mutant polypeptides. Such sequences can arise as a consequence of codon redundancy and functional equivalency that are known to occur naturally within nucleic acid sequences and the proteins thus encoded. Alternatively, functionally equivalent proteins or polypeptides can be created via the application of recombinant DNA technology, in which changes in the protein structure can be engineered, based on considerations of the properties of the amino acids being exchanged. Changes can be introduced through the application of site-directed mutagenesis techniques, e.g., to introduce improvements to the antigenicity of the protein or to test variants of a CAR or CAR LBD mutant of the present invention in order to examine the degree of lipid-binding activity, or other activity at the molecular level. Various site-directed mutagenesis techniques are known to those of skill in the art and can be employed in the present invention.

The invention further encompasses fusion proteins and peptides wherein a CAR or CAR LBD mutant coding region of the present invention is aligned within the same expression unit with other proteins or peptides having desired functions, such as for purification or immunodetection purposes.

Recombinant vectors form important further aspects of the present invention. Particularly useful vectors are those in which the coding portion of the DNA segment is positioned under the control of a promoter. The promoter can be that naturally associated with a CAR gene, as can be obtained by isolating the 5' non-coding sequences located upstream of the coding segment or exon, for example, using recombinant cloning and/or PCR technology and/or other methods known in the art, in conjunction with the compositions disclosed herein.

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In other embodiments, certain advantages can be gained by positioning the coding DNA segment under the control of a recombinant, or heterologous, promoter. As used herein, a recombinant or heterologous promoter is a promoter that is not normally associated with a CAR gene in its natural environment. Such promoters can include promoters isolated from bacterial, viral, eukaryotic, or mammalian cells. Naturally, it will be important to employ a promoter that effectively directs the expression of the DNA segment in the cell type chosen for expression. The use of promoter and cell type combinations for protein expression is generally known to those of skill in the art of molecular biology (See e.g., Sambrook & Russell, 2001, specifically incorporated herein by reference). The promoters employed can be constitutive or inducible and can be used under the appropriate conditions to direct high level expression of the introduced DNA segment, such as is advantageous in the large-scale production of recombinant proteins or peptides. One exemplary promoter system contemplated for use in high-level expression is a T7 promoter-based system.

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# XII. The Role of the Three-Dimensional Structure of the CAR LDB in Solving Additional CAR Crystals

Because polypeptides can crystallize in more than one crystal form, the structural coordinates of a CAR LBD, or portions thereof, in complex with a co-regulator as provided by the present invention, are particularly useful in solving the structure of other crystal forms of CAR and the crystalline forms of other NRs and CARs. The coordinates provided in the present invention can also be used to solve the structure of CAR or CAR LBD mutants (such as those above), CAR LDB co-complexes, or the crystalline form of any other protein with significant amino acid sequence homology to any functional domain of CAR.

One method that can be employed for the purpose of solving additional CAR crystal structures is molecular replacement. See generally, Rossmann, 1972. In the molecular replacement method, an unknown crystal form, whether it is another crystal form of a CAR or a CAR LBD, (i.e. a CAR or a

CAR LBD mutant), a CAR or a CAR LBD polypeptide in complex with another compound (i.e. a "co-complex") or the crystal of some other protein with significant amino acid sequence homology to any functional region of the CAR LBD (e.g. another NR), can be determined using the CAR LBD structure coordinates provided in Tables 2-3. This method provides an accurate structural form for the unknown crystal more quickly and efficiently than attempting to determine such information ab initio.

In addition, in accordance with this invention, CAR or CAR LBD mutants can be crystallized in complex with known modulators, such as a coregulator. The crystal structures of a series of such complexes can then be solved by molecular replacement and compared with that of wild-type CAR or the wild-type CAR LBD. Potential sites for modification within the various binding sites of the enzyme can thus be conveniently identified. This information provides an additional tool for identifying efficient binding interactions, for example, increased hydrophobic interactions between the CAR LBD and a chemical entity or compound.

All of the complexes referred to in the present disclosure can be studied using X-ray diffraction techniques (See e.g., Blundell & Johnson, 1985) and can be refined using computer software, such as the X-PLOR™ program (Brünger, 1992; X-PLOR is available from Accelrys Inc, San Diego, California, United States of America). This information can thus be used to optimize known classes of CAR and CAR LBD ligands, and more importantly, to design and synthesize novel classes of CAR and CAR LBD ligands, including co-regulators.

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#### Examples

The following Examples have been included to illustrate exemplary modes of the invention. Certain aspects of the following Examples are described in terms of techniques and procedures found or contemplated by the present inventors to work well in the practice of the invention. These Examples are exemplified through the use of standard laboratory practices of the inventors. In light of the present disclosure and the general level of skill in

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the art, those of skill will appreciate that the following Examples are intended to be exemplary only and that numerous changes, modifications, and alterations can be employed without departing from the spirit and scope of the invention.

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### Example 1

## Protein Expression and Purification

A DNA fragment encoding residues 103 - 348 of a human CAR polypeptide (GenBank Accession No. Z30425) was amplified by the polymerase chain reaction (PCR) with a commercial kit (Stratagene, La Jolla, California, United States of America). The 5' PCR primer included an Nterminal poly-histidine tag sequence (MKKGHHHHHHG; SEQ ID NO: 5) along with an Ndel endonuclease restriction site (CATATG), and the 3' PCR primer contained a BamHI restriction site (GGATCC). The PCR primers used were 5'-CGGCGCCCATATGAAAAAGGTCATCATCATCATCATCATGGTCCT GTGAACTGAGTAAGGAGCAAG-3' (SEQ ID NO: 6) and 5'-CGGCGCGCGGATCCTTAGCTGCAGATCTCCTGGAGCAGCGG 3' (SEQ ID NO: 7). The amplified DNA fragment was inserted downstream of a T7 promoter from the pRSETA vector (Invitrogen Corp., Carlsbad, California, United States of America) at the Ndel-BamHI enzyme restriction sites. E. coli cells BL21(DE3) transformed with the above expression vector were grown on a carbenicillin antibiotic agar plate (50 mg/L carbenicillin). A starter culture of 80 ml LB media (10 g/L Bacto-Tryptone, 5 g/L yeast extract, 5 g/L NaCl, QC with distilled water) with carbenicillin antibiotic (50 mg/L carbenicillin) was grown from one colony at 37°C, 250 rpm for four hours. Twelve 2 L shaker flasks with 1L LB media and carbenicillin antibiotic (50 mg/L carbenicillin) were inoculated with 5 ml of the starter culture. Cells were grown at 23°C, 250 rpm for 16 hours to an OD<sub>600</sub> of 2.0, and harvested by centrifugation. The pellet was completely resuspended with 20 ml extract buffer (150 mM NaCl. 50 mM imidazole pH 7.5) per liter of cells. The cells were sonicated for 5 minutes using a Sonicator Ultrasonic Processor XL-2015 (Heat Systems, Inc., Farmingdale, New York, United States of America) at 0°C. The lysed cells

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were centrifuged at 40,000g for 40 minutes and the supernatant was loaded on a 50 ml Ni-agarose column. The column was washed with 250 ml Buffer A (50 mM imidazole pH 7.5, 150 mM NaCl), 100 ml of Buffer B (200 mM imidazole pH 7.5, 150 mM NaCl), and the protein eluted with a 300 ml gradient to Buffer B (500 mM imidazole pH 7.5, 150 mM NaCl). The peak, which eluted at 45% Buffer B, contained 60 mg of His-tagged CAR LBD protein.

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This protein was diluted 5-fold in 10 mM Tris-Cl pH 8.0 to reduce the NaCl concentration before loading the entire sample on a 50 ml SP Sepharose FASTFLOW<sup>TM</sup> column (Pharmacia Biotech, now part of Amersham Biosciences Corp., Piscataway, New Jersey, United States of America). The column was washed with 200 ml Buffer S-A (10 mM Tris-Cl pH 8.0, 30 mM NaCl, 5 mM DTT, 1 mM EDTA pH 8.0) and the His-tagged CAR protein was eluted from the column by running a 300 ml increasing NaCl concentration gradient of Buffer S-B (10 mM Tris-Cl pH 8.0, 500 mM NaCl, 5 mM DTT, 1 mM EDTA pH 8.0). Peak fractions containing the CAR protein were pooled together, protein was concentrated to 1 mg/ml in CENTRIPREP<sup>TM</sup> 30 units (Millipore Corp., Bedford, Massachusetts, United States of America) concentrators. The protein yield was 4 mg/L cells grown. The protein was aliquoted into 10 mg aliquots at 1.0 mg/ml and stored on ice.

The purified CAR LBD protein (10 mg) was complexed with Compound 1 (10 mM in DMSO) in a 1:5 molar ratio and incubated on ice for 1 hour. The CAR LBD/Compound 1 protein complex was concentrated to 4 mg/ml in a CENTRIPREP™ 30 units and stored on ice until needed for crystallization efforts.

#### Example 2

#### Crystallization and Data Collection

CAR/Compound 1 crystals were grown at 4°C in hanging drops containing 1  $\mu$ l of the protein-ligand solutions disclosed in Example 1, and 1  $\mu$ l of well buffer (100 - 400 mM sodium potassium tartrate, pH 7.1 - 7.4). Crystals grew to a size of 100-200  $\mu$ m within several weeks. Before data

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collection, crystals were transiently mixed with the well buffer that contains an additional 14% ethylene glycol, 7% glycerol, and then flash frozen in liquid nitrogen.

Orthorhombic CAR/ligand crystals formed in the  $P2_12_12_1$  space group, with a=82.3 Å, b = 116.8 Å, c = 131.9 Å. Each asymmetric unit contained four CAR LBDs and four ligands. The crystals had a solvent content of 40%.

Crystals were screened with a Rigaku R-Axis IV detector (Rigaku International Corp., Tokyo, Japan), and data sets were collected with a MAR CCD detector at the IMCA 17ID beam line at Argonne National Labs (Argonne, Illinois, United States of America). The observed reflections were reduced, merged, and scaled with DENZO<sup>TM</sup> and SCALEPACK<sup>TM</sup> software in the HKL2000 package (Otwinowski, 1993).

## Example 3

## Structure Determination and Refinement

Structures were determined by molecular replacement methods with the CCP4 AMORE<sup>TM</sup> program (Collaborative Computational Project, 1994; Navaza, 1994) using the poly-alanine model of the conserved region of VDR LBD. Coordinates for this model are presented in Table 3.

The best fitting solution generated with the AMORE™ program gave a correlation coefficiency of 30% and an R-factor of 50%. The phases generated from molecular replacement were extensively refined and improved with solvent flattening, histogram matching, and NCS as implemented in CCP4DM and DMMULTI programs (Cowtan, 1994). Model building proceeded with QUANTA™ (available from Accelrys Inc, San Diego, California, United States of America), and refinement progressed with CNX (Brünger et al., 1998), and involved multiple cycles of manual rebuilding.

The structure of CAR in complex with the antagonist Compound 1 was determined. The statistics of the structure are summarized in Table 1.

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#### Example 4

### **Computational Analysis**

Surface area was calculated with the Connolly MS program (Connolly, 1983) and the MVP program (Lambert, 1997). The binding pocket volumes were calculated with the program GRASP (Nicholls *et al.*, 1991), using the program MVP to close openings to solvent. The sequence alignments were generated with the MVP program.

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## Example 5

## **Antagonist Assays**

Screening of synthetic compound libraries with the purified CAR LBD protein by a Fluorescence Resonance Energy Transfer (FRET) Ligand Sensing Assay (Parks et al., 1999) was conducted to identify molecules that alter the basal interaction between a coactivator peptide and the CAR LBD protein. Briefly, the purified human CAR LBD protein was biotinylated and labeled with streptavidin-conjugated fluorophore allophycocyanin. labeled CAR LBD protein was incubated with a test compound and with a peptide that included the second LXXLL binding motif of the nuclear coactivator SRC-1 (GenBank Accession No. U59302; amino acids 676-700) that was labeled with europium chelate. Data were collected with a WALLAC VICTOR<sup>TM</sup> fluorescence reader (available from PerkinElmer Life Sciences Inc., Boston, Massachusetts, United States of America) in a time resolved mode and the fluorescence ratio calculated. Compound 1 was identified from the screen to be an inverse agonist molecule that reduces the basal fluorescent signal indicating that the CAR LBD/SRC-1 interaction was reduced below background levels. Standard dose response curves were conducted with the CAR LBD protein plus Compound 1 and the EC<sub>50</sub> was determined to be 15 nM.

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#### Example 6

## Synthesis of Compound 1

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2-(benzhydrylamino) - 1 - (2-phenylethyl) - 1H - benzimidazole-6carboxamide (Compound 1) was synthesized as follows. A solution of 3fluoro-4-nitrobenzoic acid (1.28 g; 6.9 mmol) in 10 mL anhydrous N,Ndimethylformamide was treated with [O-(7-azabenzotriazol-1-yl)-1,1,3,3tetramethyluronium hexafluoro-phosphate] (2.6 g; 6.9 mmol) followed by N.N-diisopropylethylamine (3.6 ml, 20.7 mmol). After shaking for 5 min, the mixture was added to polystyrene Rink amide AM resin (1.0 g; 0.69 mmol/g; 0.69 mmol), and the reaction was rotated at 25°C for 18 h. The reaction solution was drained, and the resin was washed sequentially with N,Ndimethylformamide (3X), dichloromethane (3X), methanol (2X), and dichloromethane (3X). The dried resin was treated with 15.2 ml of a 0.5 M phenethylamine in N-methylpyrrolidinone solution then rotated at 70°C for 15 The cooled reaction was drained, and the resin was washed hours. sequentially with N,N-dimethylformamide (3X), dichloromethane (3X), methanol (2X), and dichloromethane (3X). The resin was treated with 3.8 ml of 2.0 M SnCl<sub>2</sub>-dihydrate in N-methylpyrrolidinone solution and rotated at 25 C for 24 hours. The reaction was drained and the resin washed sequentially with 30% ethylenediamine (3X)*N,N*-dimethylformamide dichloromethane (3X), methanol (2X), and dichloromethane (3X). The dried diamine resin was treated with 7.6 ml of a 0.5 M benzyhydryl isothiocyanate in N-methylpyrrolidinone solution and 7.6 ml of a 1.0 M diisopropylcarbodiimide in N-methylpyrrolidinone solution. After rotating at 80°C for 24 h the reaction was cooled to 25°C, drained, and the resin was washed sequentially with N,Ndimethylformamide (3X), dichloromethane (3X), methanol (2X), dichloromethane (3X). The resin was treated with 30 ml 95% trifluoroacetic acid (TFA) in water and rotated at 25°C for 3 hours. The resin was drained and washed with dichloromethane. The filtrate was concentrated in vacuo to give an oil. The oil was redissolved in dichloromethane and the solution was washed twice with saturated sodium bicarbonate (NaHCO<sub>3</sub>). The organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and concentrated in vacuo. The crude

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product was triturated with Et<sub>2</sub>O/hexanes, and the solid was collected by filtration to give 333 mg (98% yield) of the title compound as an off-white solid:  $^{1}$ H NMR (DMSO-d6, 400 MHz)  $\delta$  7.68 (m, 2 H), 7.63 (d, 1 H, J = 8.4), 7.54 (dd, 1 H, J = 8.0, 1.2), 7.40-7.00 (m, 17 H), 6.36 (d, 1 H, J = 8), 4.42 (t, 2 H, J = 7.4), 2.97 (t, 2 H, J = 7.4); MS (ESP+) m/e 447 (MH<sup>+</sup>).

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Table 2

Atomic Structure Coordinate Data Obtained From

X-ray Diffraction From the Ligand-binding Domain of CAR

In Complex With Compound 1

	ATOM	1	N	LEU A	A	120	34.417	18.787	67.312	1.00	50.31	N
	MOTA	2	CA	LEU A	A	120	34.298	17.304	67.212	1.00	49.96	С
	ATOM	3	С	LEU A	Α	120	33.672	16.891	65.886	1.00	49.44	С
15	ATOM	4	0	LEU /	Α	120	32.815	17.592	65.344	1.00	49.49	0
	MOTA	5	CB	LEU A	Α	120	33.447	16.756	68.363		50.64	C
	MOTA	6	CG	LEU .	A	120	34.003	16.880	69.783	1.00	51.38	C
	MOTA	7	CD1	LEU .	Α	120	32.969	16.374	70.777	1.00	51.56	C
	MOTA	8	CD2	LEU .	Α	120	35.297	16.085	69.906	1.00	51.43	С
20	MOTA	9	N	ARG .	A	121	34.106	15.745	65.375	1.00	48.14	N
	MOTA	10	CA	ARG	A	121	33.599	15.221	64.117	1.00	47.01	С
	MOTA	11	С	ARG	Α	121	33.113	13.790	64.314	1.00	45.50	С
	MOTA	12	0	ARG			33.775	12.836	63.905	1.00	45.36	0
	MOTA	13	CB	ARG	Α	121	34.700	15.264	63.052	1.00	48.45	C
25	ATOM	14	CG	ARG			35.233	16.664	62.790	1.00	49.89	С
	ATOM	15	CD	ARG			36.430	16.655	61.852	1.00	52.32	С
	ATOM	16	NE	ARG			36.100	16.133	60.529	1.00	53.49	N
	MOTA	17	CZ	ARG			36.947	16.112	59.504	1.00	54.08	С
	MOTA	18	NH1	_		_	38.178	16.586	59.648	1.00	54.50	N
30	MOTA	19	NH2	ARG			36.563	15.620	58.334		54.12	N
	MOTA	20	N	PRO			31.946	13.622	64.955		43.87	N
	ATOM	21	CA	PRO			31.403	12.282	65.187		42.99	С
	ATOM	22	C	PRO			31.173	11.529	63.881		42.25	C
	ATOM	23	0	PRO			30.823	12.125	62.862		42.01	0
35	MOTA	24	CB	PRO			30.105	12.561	65.944	1.00		С
	MOTA	25	CG			122	29.699	13.908	65.437	1.00	43.60	С
	ATOM	26	CD			122	31.010	14.655	65.429	1.00	43.27	С
	MOTA	27	N	LYS			31.379	10.218	63.920	1.00		N
	ATOM	28	CA	LYS			31.205	9.378	62.744	1.00	41.30	С
40	ATOM	29	С			123	29.732	9.158	62.431		40.35	С
	ATOM	30	0			123	28.877	9.250	63.313	1.00		0
	MOTA	31	CB	LYS			31.885	8.024	62.965	1.00		С
	MOTA	32	CG			123	33.371	8.127	63.279	1.00		С
	MOTA	33	CD			123	33.979	6.761	63.564	1.00		С
45	MOTA	34	CE			123	35.463	6.876	63.882	1.00		С
	MOTA	35	NZ			123	36.066	5.558	64.225	1.00		N
	ATOM	36	N			124	29.439	8.879	61.165	1.00		N
	ATOM	37	CA			124	28.071	8.622	60.744	1.00		С
	MOTA	38	С	LEU	A	124	27.606	7.325	61.384	1.00	38.41	С

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	MOTA	39		LEU A		28.293	6.308	61.304		2
	MOTA	40		LEU A		27.996	8.491	59.220		2
	MOTA	41		LEU A		28.162	9.776	58.406		2
_	ATOM	42		LEU A		28.401	9.438	56.941		2
5	MOTA	43		LEU A		26.922	10.633	58.564		С
	ATOM	44	N	SER A		26.448	7.362	62.029		N
	MOTA	45	CA	SER A		25.905	6.168	62.661		C
	MOTA	46	С	SER A		25.496	5.197	61.561		С
	MOTA	47	0	SER A		25.386	5.581	60.395		0
10	MOTA	48	CB	SER A		24.679	6.523	63.495		C
	MOTA	49	OG	SER A		23.619	6.951	62.660		0
	MOTA	50	N	GLU A		25.271	3.940	61.923		N
	MOTA	51	CA	GLU A		24.865	2.956	60.930		С
	MOTA	52	С	GLU A		23.535	3.385	60.314		C
15	MOTA	53	0	GLU A		23.313	3.207	59.115		0
	ATOM	54	CB	GLU A	126	24.727	1.573	61.573		С
	MOTA	55	CG	GLU A	126	24.325	0.463	60.605		С
	MOTA	56	CD	GLU A	126	25.202	0.414	59.361	1.00 51.93	С
	MOTA	57	OE1	GLU A	126	24.878	1.105	58.366	1.00 53.34	0
20	ATOM	58	OE2	GLU A	126	26.222	-0.308	59.379	1.00 53.64	0
	MOTA	59	N	GLU A	127	22.659	3.960	61.133	1.00 40.27	N
	MOTA	60	CA	GLU A	127	21.358	4.412	60.650	1.00 39.52	C
	MOTA	61	C	GLU A	127	21.512	5.550	59.647	1.00 37.38	С
	ATOM	62	0	GLU A	127	20.814	5.594	58.630	1.00 36.24	0
25	MOTA	63	CB	GLU A	127	20.481	4.891	61.807	1.00 41.53	С
	ATOM	64	CG	GLU A	127	19.091	5.320	61.363	1.00 45.78	С
	ATOM	65	CD	GLU A	127	18.236	5.832	62.504	1.00 47.87	С
	ATOM	66	OE1			18.572	6.890	63.075	1.00 49.93	0
	ATOM	67	OE2	GLU A	127	17.227	5.173	62.832	1.00 50.45	0
30	ATOM	68	N	GLN A	128	22.420	6.473	59.939	1.00 34.92	N
	ATOM	69	CA	GLN A	128	22.654	7.603	59.052	1.00 33.94	C
	ATOM	70	С	GLN A	128	23.239	7.134	57.721	1.00 34.19	С
	ATOM	71	0	GLN A		22.905	7.671	56.665	1.00 32.45	0
	ATOM	72	CB	GLN A		23.573	8.622	59.735	1.00 33.20	С
35	ATOM	73	CG	GLN A		22.861	9.410	60.835	1.00 32.00	С
•	ATOM	74	CD	GLN A		23.785	10.317	61.629	1.00 32.20	С
	ATOM	75	OE1			23.346	11.326	62.192	1.00 33.66	0
	ATOM	76	NE2			25.061	9.960	61.691	1.00 30.80	N
	ATOM	77	N	GLN A		24.101	6.124	57.768	1.00 33.75	N
40	ATOM	78	CA	GLN A		24.692	5.591	56.545	1.00 35.00	C
	ATOM	79	C	GLN A		23.588	4.965	55.702	1.00 34.31	C
	ATOM	80		GLN A		23.562	5.111			Ō
	ATOM	81	СВ	GLN A		25.747	4.531	56.874	1.00 37.89	C
	ATOM	82	CG	GLN A		26.977	5.078	57.579	1.00 42.41	Č
45	ATOM	83	CD	GLN A		27.983	3.995	57.929	1.00 45.15	Č
40	ATOM	84	OE1			28.998	4.261	58.575	1.00 46.46	ō
	ATOM	85	NE2			27.704	2.766	57.504	1.00 46.27	N
	ATOM	86	N	ARG A		22.674	4.270	56.370	1.00 33.44	N
	ATOM	87	CA	ARG A		21.556	3.614	55.703	1.00 34.05	C
50	ATOM	88	C	ARG A		20.653	4.638	55.018	1.00 32.98	č
30		89	ŏ	ARG A		20.226	4.436	53.881	1.00 31.44	ŏ
	MOTA ATOM	90	СВ	ARG A			2.794	56.723	1.00 37.04	C
	MOTA		CG	ARG A			2.794	56.723	1.00 37.04	C
	MOTA	91 92	CD	ARG A				57.171	1.00 41.38	C
55	MOTA						1.108	56.790		N
55	MOTA	93	NE	ARG A			0.608			
	ATOM	94		ARG A			1.291	56.938		C
	MOTA	95		L ARG A				57.465		N
	MOTA	96	NH	2 ARG A	1 T20	15.357	0.757	56.556	1.00 52.73	N

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	MOTA	97	N	ILE A	A 1	.31	20.367	5.735	55.712	1.00	31.16	N
	ATOM	98	CA	ILE A	A 1	131	19.519	6.790	55.158	1.00	30.41	C
	ATOM	99	С	ILE A	A 1	131	20.120	7.343	53.865	1.00	29.21	С
	MOTA	100	0	ILE A	A 1	L31	19.414	7.528	52.872	1.00	27.86	0
5	ATOM	101	СВ	ILE A	A 1	131	19.334	7.945	56.177	1.00	31.61	C
	ATOM	102	CG1	ILE A	A 1	131	18.513	7.448	57.372	1.00	32.47	C
	ATOM	103	CG2	ILE 2	A 1	131	18.657	9.138	55.507	1.00	31.13	С
	ATOM	104		ILE 2			18.287	8.496	58.457	1.00	33.63	С
	ATOM	105	N	ILE			21.424	7.601	53.876		28.81	Ŋ
10	ATOM	106	CA	ILE .			22.094	8.124	52.691		29.13	С
	ATOM	107	С	ILE .			22.029	7.115	51.544	1.00	29.37	C
	ATOM	108	0	ILE			21.786	7.486	50.394		28.72	Ō
	ATOM	109	СВ	ILE .			23.570	8.468	52.994		29.90	C
	ATOM	110	CG1	ILE .			23.628	9.625	53.995	•	30.31	C
15	ATOM	111	CG2	ILE			24.306	8.838	51.708		30.32	Č
	ATOM	112	CD1				25.027	9.997	54.432		31.33	Č
	ATOM	113	N	ALA			22.239	5.841	51.862		28.31	N
	ATOM	114	CA	ALA			22.203	4.785	50.851		27.51	C
	ATOM	115	C	ALA			20.820	4.680	50.213		26.94	Č
20	ATOM	116	Ö	ALA			20.694	4.542	48.993		26.91	ō
	ATOM	117	СВ	ALA			22.587	3.454	51.479		27.94	Č
	ATOM	118	N	ILE			19.786	4.739	51.044		26.00	N
	ATOM	119	CA	ILE			18.413	4.659	50.564		25.19	C
	ATOM	120	C	ILE			18.090	5.832	49.643		24.84	č
25	ATOM	121	ō	ILE			17.490	5.651	48.585		23.10	ō
	ATOM	122	СВ	ILE			17.416	4.660	51.742		26.47	č
	ATOM	123	CG1				17.511	3.331	52.493		27.92	Č
	ATOM	124	CG2				15.997	4.901	51.239		26.56	c
	ATOM	125	CD1				16.714	3.297	53.778		29.71	Č
30	MOTA	126	N	LEU			18.494	7.030	50.047		23.54	N
00	ATOM	127	CA	LEU			18.228	8.220	49.242		23.28	C
	MOTA	128	C	LEU			18.987	8.217	47.914		22.05	C
	MOTA	129	Ö	LEU			18.454	8.656	46.894		21.44	Ö
	MOTA	130	СВ	LEU			18.559	9.480	50.045		23.21	C
35	ATOM	131	CG	LEU			17.644	9.754	51.246		24.57	C
33	ATOM	132		LEU			18.057	11.076	51.900		26.44	C
	ATOM	133		LEU			16.185	9.820	50.789		25.56	C
	ATOM	134	N N	LEU			20.223	7.725	47.913		22.40	N
	ATOM	135	CA	LEU			20.223	7.675	46.669		23.29	C
40		136	C	LEU			20.302	6.721		1.00		C
40	ATOM ATOM	137	0	LEU			20.302	6.721	45.705 44.512		23.30	0
		138	СВ	LEU			22.424	7.194	46.920		24.60	C
	ATOM		CG	LEU			23.395	8.196	47.549		25.56	
	ATOM	139 140		LEU			24.740	7.518	47.798		26.67	C
45	ATOM	141		LEU			23.555	9.398	46.628		26.04	C
40	ATOM	142	N N	ASP				5.591	46.232		23.87	C
	MOTA		CA	ASP			19.845					N
	ATOM	143					19.156	4.589	45.427		23.95	C
	MOTA	144	C	ASP			17.844	5.152	44.870		23.67	C
EΩ	MOTA	145	0	ASP			17.513	4.943	43.697			0
50	MOTA	146	CB	ASP			18.886	3.348	46.282		26.93	C
	MOTA	147	CG			137	18.158	2.266	45.524		31.10	C
	ATOM	148		ASP			17.010	1.947	45.900		34.78	
	ATOM	149		ASP			18.730	1.734	44.552		34.13	
EE	ATOM	150	N			138	17.105	5.867	45.714		22.31	
55	ATOM	151	CA			138		6.472	45.312		22.31	C
	ATOM	152	C			138		7.435	44.157		21.39	
	ATOM	153	0			138		7.445	43.183		20.83	
	ATOM	154	CB	ALA	Α	138	15.213	7.219	46.487	1.00	23.04	С

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	MOTA	155	N	HIS A	139	17.107	8.249	44.263	1.00 21.06	N
	MOTA	156	CA	HIS A	139	17.408	9.202	43.208	1.00 21.28	С
	MOTA	157	С	HIS A	139	17.814	8.511	41.905	1.00 21.64	С
	MOTA	158	0	HIS A	139	17.385	8.913	40.824	1.00 21.17	0
5	MOTA	159		HIS A		18.528	10.152	43.631	1.00 21.21	С
	MOTA	160	CG	HIS A	139	18.730	11.288	42.680	1.00 22.53	С
	MOTA	161		HIS A		19.955	11.593	42.126	1.00 25.49	N
	MOTA	162	CD2	HIS A	139	17.850	12.173	42.157	1.00 19.49	С
	MOTA	163	CE1	HIS F	139	19.820	12.615	41.300	1.00 20.82	С
10	MOTA	164	NE2	HIS A	139	18.552	12.986	41.301	1.00 23.99	N
	MOTA	165	N	HIS A	140	18.650	7.479	42.005	1.00 21.50	N
	MOTA	166	CA	HIS A	140	19.099	6.760	40.819	1.00 22.20	С
	MOTA	167	С	HIS A	140	17.947	6.088	40.082	1.00 21.95	С
	ATOM	168	0	HIS A	140	17.997	5.911	38.861	1.00 21.87	0
15	MOTA	169	CB	HIS A	A 140	20.153	5.710	41.193	1.00 23.76	С
	ATOM	170	CG	HIS A	A 140	21.398	6.291	41.787	1.00 25.80	С
	ATOM	171	ND1	HIS A	A 140	21.803	7.585	41.546	1.00 27.26	N
	ATOM	172	CD2	HIS A	A 140	22.341	5.745	42.591	1.00 26.22	С
	MOTA	173	CE1	HIS A	A 140	22.942	7.814	42.176	1.00 26.08	С
20	MOTA	174	NE2	HIS A	A 140	23.291	6.714	42.817	1.00 27.71	N
	MOTA	175	N	LYS 2	A 141	16.908	5.719	40.821	1.00 20.41	N
	MOTA	176	CA		A 141	15.745	5.071	40.225	1.00 21.89	С
	MOTA	177	С	LYS	A 141	14.746	6.078	39.665	1.00 21.31	C
	ATOM	178	0	LYS	A 141	13.916	5.730	38.832	1.00 22.47	0
25	ATOM	179	СВ		A 141	15.031	4.203	41.265	1.00 23.28	C
	ATOM	180	CG		A 141	15.804	2.960	41.668	1.00 26.83	С
	ATOM	181	CD		A 141	15.080	2.209	42.771	1.00 30.63	С
	ATOM	182	CE	LYS	A 141	15.781	0.902	43.093	1.00 33.64	С
	ATOM	183	NZ	LYS	A 141	15.122	0.206	44.231	1.00 36.58	N
30	MOTA	184	N		A 142	14.840	7.325	40.107	1.00 20.65	N
-	ATOM	185	CA		A 142	13.893	8.348	39.664	1.00 20.68	C
	ATOM	186	С		A 142	14.440	9.502	38.833	1.00 20.45	С
	ATOM	187	0		A 142	13.682	10.375	38.420	1.00 20.32	0
	ATOM	188	СВ	THR	A 142	13.142	8.935	40.865	1.00 20.48	C
35	MOTA	189	OG1		A 142	14.081	9.474	41.805	1.00 18.91	0
••	ATOM	190	CG2		A 142	12.326	7.850	41.546	1.00 19.94	C
	ATOM	191	N		A 143	15.747	9.520	38.595	1.00 20.03	N
	ATOM	192	CA		A 143	16.342	10.566	37.768	1.00 20.44	С
	ATOM	193	С		A 143	17.207	9.895	36.706	1.00 20.75	C
40	ATOM	194	0		A 143	18.248	9.323	37.013	1.00 21.56	0
	ATOM	195	СВ		A 143	17.198	11.529	38.610	1.00 20.88	С
	ATOM	196			A 143	17.673	12.742			С
	ATOM	197			A 143	18.721	12.650	36.915	1.00 21.44	С
	MOTA	198			A 143		13.980	37.994	1.00 21.13	C
45	ATOM	199			A 143		13.762	36.170	1.00 21.80	C
	ATOM	200			A 143		15.090	37.253	1.00 20.26	C
	ATOM	201	CZ		A 143		14.978	36.347	1.00 22.15	C
	ATOM	202	ОН		A 143		16.077	35.612	1.00 21.28	0
	ATOM	203	N		A 144		9.959	35.461	1.00 20.48	N
50	ATOM	204	CA		A 144		9.365	34.326	1.00 21.36	C
	ATOM	205	C		A 144		10.387	33.751	1.00 22.06	Č
	ATOM	206	ō		A 144		11.348	33.102	1.00 21.75	Ö
	ATOM	207	СВ		A 144		8.955	33.274	1.00 21.65	Č
	ATOM	208	CG		A 144		8.481	31.976	1.00 22.22	č
55	MOTA	209			A 144		8.286	31.921	1.00 22.12	Ö
55	ATOM	210			A 144			31.007	1.00 23.20	ŏ
	MOTA	211	N		A 145			33.976	1.00 21.93	N
	ATOM	212	CA		A 145			33.483		C
	AION	212	J.,	- 110	1-30	20.113	*****	55.405	1.00 25.05	-

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	MOTA	213	С	PRO A	145	20.968	11.106	31.968	1.00 22.50	С
	ATOM	214	0	PRO A	145	21.754	11.906	31.451	1.00 23.61	0
	MOTA	215	CB	PRO A	A 145	22.026	10.620	34.225	1.00 23.45	С
	ATOM	216	CG	PRO A	A 145	21.809	9.150	34.297	1.00 24.95	С
5	MOTA	217	CD	PRO A	A 145	20.347	9.052	34.700	1.00 23.26	С
	MOTA	218	N	THR A	A 146	20.265	10.224	31.256	1.00 22.03	N
	MOTA	219	CA	THR A	A 146	20.364	10.192	29.796	1.00 21.95	С
	MOTA	220	С	THR A	A 146	19.174	10.907	29.155	1.00 22.52	С
	MOTA	221	0	THR A	A 146	19.181	11.177	27.953	1.00 22.17	0
10	MOTA	222	CB	THR A	A 146	20.433	8.750	29.233	1.00 21.96	С
	MOTA	223	OG1	THR A	A 146	19.167	8.099	29.395	1.00 21.08	0
	MOTA	224	CG2	THR Z	A 146	21.509	7.949	29.956	1.00 23.14	C
	MOTA	225	N	TYR A	A 147	18.158	11.210	29.963	1.00 22.04	N
	MOTA	226	CA	TYR 2	A 147	16.963	11.912	29.489	1.00 22.53	C
15	ATOM	227	С	TYR 2	A 147	16.313	11.191	28.309	1.00 23.10	С
	MOTA	228	0	TYR .	A 147	15.789	11.821	27.393	1.00 23.05	0
	ATOM	229	СВ	TYR .	A 147	17.335	13.350	29.093	1.00 23.34	С
	MOTA	230	CG	TYR .	A 147	18.159	14.049	30.150	1.00 23.73	С
	ATOM	231	CD1	TYR .	A 147	19.525	14.274	29.968	1.00 25.15	С
20	MOTA	232	CD2	TYR .	A 147	17.593	14.398	31.372	1.00 23.61	C
	ATOM	233	CE1	TYR	A 147	20.304	14.818	30.989	1.00 25.82	,C
	ATOM	234	CE2	TYR	A 147	18.363	14.941	32.396	1.00 26.56	С
	ATOM	235	CZ		A 147	19.716	15.142	32.199	1.00 26.11	C
	ATOM	236	ОН		A 147	20.484	15.619	33.237	1.00 29.64	0
25	MOTA	237	N		A 148	16.326	9.862	28.355	1.00 23.29	N
	ATOM	238	CA		A 148	15.781	9.046	27.278	1.00 23.65	C
	ATOM	239	C		A 148	14.263	9.078	27.073	1.00 24.65	Č
	ATOM	240	ō		A 148	13.783	8.650	26.024	1.00 24.62	Ō
	ATOM	241	СВ		A 148	16.243	7.593	27.450	1.00 26.66	C
30	ATOM	242	OG		A 148	15.684	7.006	28.614	1.00 29.82	ō
00	ATOM	243	N		A 149	13.505	9.576	28.048	1.00 22.99	N
	ATOM	244	CA		A 149	12.045	9.632	27.905	1.00 23.85	C
	ATOM	245	C		A 149	11.534	10.925	27.272	1.00 24.00	Č
	ATOM	246	Ö		A 149	10.371	11.008	26.879	1.00 24.41	ō
35	ATOM	247	СВ		A 149	11.349	9.488	29.263	1.00 24.47	Č
55	ATOM	248	CG		A 149	11.517	8.114	29.872	1.00 27.05	C
	ATOM	249			A 149	11.441	7.116	29.124	1.00 26.86	Ö
	ATOM	250	OD2		A 149	11.707	8.037	31.105	1.00 26.29	o
	ATOM	251	N		A 150	12.396	11.927	27.171	1.00 24.31	N
40	MOTA	252	CA		A 150	11.995	13.231	26.646	1.00 25.09	C
70	ATOM	253	C		A 150	11.363	13.263	25.252	1.00 25.91	C
	MOTA	254	Õ		A 150			24.949		
	ATOM	255	СВ		A 150		14.187	26.715	1.00 24.68	C
	ATOM	256	CG		A 150		14.611	28.121	1.00 25.17	C
45	ATOM	257			A 150		13.726	29.187	1.00 25.54	C
70	ATOM	258			A 150		15.720	28.374	1.00 26.43	C
		259			A 150		14.104	30.484	1.00 25.74	c
	MOTA	260			A 150			29.667	1.00 25.74	c
	MOTA		CZ		A 150		15.386		1.00 24.63	C
50	ATOM	261 262	N		A 151			30.721 24.404		N
30	MOTA									
	MOTA	263	CA C		A 151 A 151			23.063 23.094	1.00 28.74 1.00 28.90	C
	MOTA	264								
	MOTA	265	O		A 151			22.075		0
EE	ATOM	266	CB		A 151			22.154		C
55	MOTA	267	SG		A 151			22.716		
	ATOM	268	N		A 152					
	MOTA	269	CA		A 152					
	MOTA	270	С	GLN	A 152	6.858	12.285	24.711	1.00 27.73	С

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	ATOM	271	0	GLN A 152	5.633	12.202	24.590	1.00 28.51	0
	ATOM	272	CB	GLN A 152	7.602	10.021	25.473	1.00 29.61	Ċ
	ATOM	273	CG	GLN A 152	8.312	8.724	25.123	1.00 33.35	C
	ATOM	274	CD	GLN A 152	8.121	7.650	26.173	1.00 36.62	C
5	ATOM	275		GLN A 152	6.995	7.260	26.478	1.00 39.37	Ō
_	MOTA	276	NE2		9.225	7.162	26.732	1.00 38.35	N
	ATOM	277	N	PHE A 153	7.469	13.395	25.115	1.00 25.45	N
	MOTA	278	CA	PHE A 153	6.705	14.597	25.439	1.00 25.30	C
	ATOM	279	C	PHE A 153	6.261	15.273	24.151	1.00 25.61	Č
10	ATOM	280	Ö	PHE A 153	6.799	14.998	23.071	1.00 24.69	Ö
. •	ATOM	281	СВ	PHE A 153	7.564	15.608	26.215	1.00 23.94	c
	MOTA	282	CG	PHE A 153	8.187	15.060	27.469	1.00 23.45	c
	ATOM	283		PHE A 153	9.332	15.654	27.990	1.00 23.45	C
	MOTA	284		PHE A 153	7.654	13.034	28.116	1.00 22.73	c
15	ATOM	285	CE1	PHE A 153	9.948	15.146	29.133	1.00 23.40	c
15	ATOM	286	CE2	PHE A 153	8.261	13.434	29.133		C
	ATOM	287	CZ	PHE A 153				1.00 22.50	
					9.414	14.037	29.769	1.00 22.91	C
	MOTA	288	N	ARG A 154	5.276	16.158	24.260	1.00 25.51	
20	MOTA	289	CA	ARG A 154	4.842	16.902	23.092	1.00 26.08	
20	MOTA	290	C	ARG A 154	6.094	17.673	22.689	1.00 27.20	-
	MOTA	291	0	ARG A 154	6.824	18.184	23.542	1.00 26.99	
	MOTA	292	СВ	ARG A 154	3.681	17.830	23.449	1.00 26.73	
	MOTA	293	CG	ARG A 154	2.351	17.087	23.522	1.00 27.85	
<b>^</b> =	MOTA	294	CD	ARG A 154	1.232	17.964	24.066	1.00 27.71	
25	MOTA	295	NE	ARG A 154	1.347	18.138	25.509	1.00 27.14	
	MOTA	296	CZ	ARG A 154	0.497	18.839	26.248	1.00 28.47	С
	MOTA	297	NH1		-0.538	19.444	25.677	1.00 29.16	N
	MOTA	298	NH2	ARG A 154	0.673	18.919	27.560	1.00 27.66	N
	ATOM	299	N	PRO A 155	6.368	17.757	21.384	1.00 27.28	N
30	MOTA	300	CA	PRO A 155	7.554	18.454	20.892	1.00 28.12	С
	MOTA	301	С	PRO A 155	7.709	19.929	21.217	1.00 28.41	C
	MOTA	302	0	PRO A 155	6.733	20.676	21.291	1.00 27.77	0
	MOTA	303	CB	PRO A 155	7.491	18.206	19.388	1.00 28.83	С
	MOTA	304	CG	PRO A 155	6.020	18.191	19.130	1.00 29.19	C
35	MOTA	305	CD	PRO A 155	5.508	17.335	20.262	1.00 28.61	
	MOTA	306	N	PRO A 156	8.956	20.361	21.437	1.00 28.25	N
	MOTA	307	CA	PRO A 156	9.202	21.768	21.739	1.00 29.56	
	ATOM	308	С	PRO A 156	9.054	22.532	20.425	1.00 30.08	
	ATOM	309	0	PRO A 156	9.483	22.054	19.371	1.00 30.96	
40	MOTA	310	CB	PRO A 156	10.640	21.763	22.250	1.00 29.92	
	MOTA	311	CG	PRO A 156	11.262	20.646	21.476	1.00 30.45	_
	ATOM	312	CD	PRO A 156	10.198	19.573	21.538	1.00 29.15	
	ATOM	313	N	VAL A 157	8.417	23.693	20.489	1.00 30.75	-
	ATOM	314	CA	VAL A 157	8.220	24.538	19.319	1.00 31.52	
45	ATOM	315	C	VAL A 157	8.764	25.907	19.692	1.00 32.33	
	ATOM	316	ŏ	VAL A 157	8.361	26.482	20.698	1.00 33.09	
	ATOM	317	СВ	VAL A 157	6.727	24.663	18.962	1.00 33.03	
	ATOM	318		VAL A 157	6.544	25.654	17.825	1.00 32.48	
		319		VAL A 157	6.177	23.302			
50	MOTA						18.573	1.00 32.24	
50	MOTA	320	N	ARG A 158	9.681	26.425	18.885	1.00 33.83	
	ATOM	321	CA	ARG A 158	10.289	27.716	19.173	1.00 36.19	
	ATOM	322	С	ARG A 158	10.020	28.766	18.096	1.00 38.44	
	ATOM	323	0	ARG A 158	10.763	28.881	17.123	1.00 39.20	
EE	ATOM	324	CB	ARG A 158	11.794	27.523	19.367	1.00 35.80	
55	MOTA	325	CG	ARG A 158	12.131	26.585	20.524	1.00 34.7	
	ATOM	326	CD	ARG A 158	13.606	26.231	20.561	1.00 35.0	
	MOTA	327	NE	ARG A 158	13.991	25.641	21.841	1.00 32.63	
	MOTA	328	CZ	ARG A 158	14.006	24.339	22.113	1.00 31.8	2 C

	ATOM	329	NH1	ARG Z	A 1	58	13.658	23.450	21.192	1.00	32.10	N
	MOTA	330	NH2	ARG A	A 1	58	14.370	23.926	23.319	1.00	29.69	N
	ATOM	331	N	VAL 2	A 1	59	8.949	29.531	18.284	1.00	40.67	N
	ATOM	332	CA	VAL 2	A 1	59	8.568	30.574	17.338	1.00	42.44	С
5	MOTA	333	С	VAL 2	A 1	59	9.511	31.767	17.432	1.00	43.24	С
	ATOM	334	0	VAL 2	A 1	59	10.170	31.968	18.451	1.00	42.85	0
	MOTA	335	CB	VAL 2	A 1	59	7.135	31.066	17.607	1.00	42.85	С
	ATOM	336	CG1	VAL .	A 1	59	6.147	29.937	17.367	1.00	43.48	С
	ATOM	337	CG2	VAL .	A 1	59	7.027	31.577	19.040	1.00	43.60	С
10	MOTA	338	N	ASN .	A 1	60	9.576	32.557	16.365	1.00	44.06	N
	MOTA	339	CA	ASN .	A 1	60	10.440	33.730	16.357	1.00	44.92	С
	MOTA	340	С	ASN .	A 1	60	9.876	34.768	17.320	1.00	45.24	С
	MOTA	341	0	ASN .	A 1	60	8.728	35.198	17.185	1.00	45.27	0
	MOTA	342	CB	ASN .	A 1	60	10.530	34.326	14.949	1.00	46.00	С
15	MOTA	343	CG	ASN	A 1	60	11.017	33.322	13.921	1.00	47.25	С
	MOTA	344	OD1	ASN	A 1	60	12.030	32.649	14.124	1.00	47.25	0
	ATOM	345	ND2	ASN	A 1	60	10.298	33.218	12.808		48.36	N
	ATOM	346	N	ASP	A 1	61	10.688	35.156	18.298		45.02	N
	ATOM	347	CA	ASP	A 1	61	10.282	36.142	19.289	1.00	44.79	С
20	MOTA	348	С	ASP	A 1	61	11.515	36.834	19.862	1.00	44.74	С
	ATOM	349	0	ASP	A 1	61	11.679	36.939	21.077	1.00	44.64	0
	MOTA	350	СВ	ASP			9.483	35.463	20.406		44.26	C
	MOTA	351	CG	ASP	A 1	61	9.101	36.421	21.515	1.00	44.34	С
	MOTA	352	OD1	ASP	A 1	61	8.640	37.540	21.201	1.00	43.26	0
25	ATOM	353	OD2	ASP	A 1	61	9.258	36.054	22.700	1.00	43.90	0
	ATOM	354	N	GLY	A 1	62	12.383	37.304	18.972		44.73	N
	ATOM	355	CA	GLY	A 1	62	13.592	37.977	19.409		44.74	С
	MOTA	356	С	GLY	A 1	62	13.292	39.196	20.261		44.56	С
	MOTA	357	0	GLY			14.135	39.638	21.042	1.00	45.10	0
30	ATOM	358	N	GLY			12.086	39.736	20.116		44.30	N
	MOTA	359	CA	GLY	A 1	.63	11.706	40.911	20.879		43.74	C
	MOTA	360	С	GLY	A 1	.63	11.206	40.618	22.282		43.23	С
	MOTA	361	0	GLY			11.066	41.533	23.096		43.53	0
	ATOM	362	N	GLY			10.946	39.346	22.572		42.43	N
35	MOTA	363	CA	GLY			10.450	38.980	23.889		40.70	С
	MOTA	364	С	GLY			9.094	39.616	24.130		39.47	C
	MOTA	365	0	GLY	A 1	64	8.812	40.125	25.222	1.00	40.10	0
	ATOM	366	N	SER	A 2	216	8.256	39.587	23.099		36.82	N
	MOTA	367	CA	SER	A 2	216	6.918	40.165	23.162	1.00	35.37	C
40	ATOM	368	С	SER	A 2	216	5.965	39.359	24.032	1.00	34.15	С
	MOTA	369	0	SER	A 2	216	5.653	38.213	23.721	1.00	32.50	0
	ATOM	370	CB	SER	A 2	216	6.329	40.277	21.755		35.39	С
	ATOM	371	OG	SER	A 2	216	4.958	40.634	21.812	1.00	35.41	0
	ATOM	372	N	VAL	A 2	217	5.495	39.969	25.116		33.39	N
45	ATOM	373	CA	VAL	A 2	217	4.563	39.301	26.013		33.22	С
	MOTA	374	С	VAL			3.299	38.922	25.251		32.19	С
	ATOM	375	0	VAL	A 2	217	2.783	37.816	25.399		31.92	0
	ATOM	376	СВ	VAL	A 2	217	4.161	40.208	27.195		33.21	С
	ATOM	377	CG1	. VAL	A 2	217	3.203	39.462	28.119		35.52	С
50	MOTA	378	CG2	VAL	A :	217	5.396	40.644	27.960		35.70	С
	MOTA	379	N	THR	A :	218	2.809	39.846	24.428		31.30	N
	MOTA	380	CA	THR	<b>A</b> :	218	1.597	39.609	23.653		30.58	C
	MOTA	381	С	THR			1.736	38.398	22.741		30.30	C
	ATOM	382	0	THR			0.852	37.544	22.695		30.29	ō
55	ATOM	383	CB	THR			1.235	40.843	22.802		30.65	Ċ
	ATOM	384		THR			1.025	41.966	23.667		30.30	ō
	ATOM	385		? THR			-0.035	40.587	22.000		31.23	Č
	ATOM	386	N	LEU			2.849	38.325	22.018		29.44	N
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	ATOM	387	CA	LEU A	219	3.095	37.206	21.117	1.00 29.87	
	ATOM	388	С	LEU A		3.260	35.905	21.894	1.00 29.21	
	MOTA	389	0	LEU A		2.710	34.869	21.516	1.00 29.73	
_	MOTA	390	CB	LEU A		4.355	37.462	20.286	1.00 31.48	
5	MOTA	391	CG	LEU A		4.778	36.321	19.352	1.00 33.59	
	MOTA	392		LEU A		3.700	36.083	18.301	1.00 34.93	
	MOTA	393		LEU F		6.100	36.676	18.690	1.00 35.5	
	MOTA	394	N	GLU A		4.018	35.963	22.982	1.00 28.83	
40	ATOM	395	CA	GLU A		4.258	34.781	23.801	1.00 29.0	
10	ATOM	396	C	GLU A		2.958	34.194	24.342	1.00 29.0	
	MOTA	397	0	GLU A		2.757	32.983	24.297	1.00 27.8	_
	MOTA	398	CB	GLU A		5.213	35.131	24.946	1.00 31.3	
	MOTA	399	CG	GLU A		6.620	35.466	24.456	1.00 32.7	_
4 =	MOTA	400	CD	GLU A		7.434	36.277	25.450	1.00 35.6	
15	ATOM	401	OE1			8.574	36.657	25.104	1.00 37.0	
	ATOM	402		GLU A		6.944	36.541	26.569	1.00 36.2	
	ATOM	403	N		A 221	2.073	35.052	24.841	1.00 28.7	
	ATOM	404	CA		A 221 A 221	0.799	34.592	25.383	1.00 29.8	
20	ATOM	405	C			-0.143 -0.923	34.089	24.293	1.00 29.7 1.00 30.0	
20	MOTA	406 407	O CB		A 221 A 221	0.125	33.165 35.714	24.516 26.181	1.00 30.0	
	MOTA MOTA	407	CG		A 221	0.743	36.046	27.544	1.00 30.0	
	MOTA	409		LEU		0.743	37.278	28.138	1.00 31.0	
	ATOM	410		LEU		0.588	34.850	28.482	1.00 32.2	
25	MOTA	411	N N		A 222	-0.066	34.687	23.108	1.00 31.0	
25	ATOM	412	CA		A 222	-0.931	34.272	22.011	1.00 32.2	
	ATOM	413	C		A 222	-0.536	32.905	21.460	1.00 32.8	
	ATOM	414	ŏ		A 222	-1.380	32.170	20.947	1.00 33.7	
	ATOM	415	СВ		A 222	-0.895	35.304	20.877	1.00 34.8	
30	ATOM	416	OG		A 222	0.367	35.315	20.230	1.00 39.0	
00	ATOM	417	N		A 223	0.742	32.558	21.584	1.00 31.8	
	ATOM	418	CA		A 223	1.234	31.288	21.063	1.00 31.7	
	ATOM	419	C		A 223	1.596	30.215	22.089	1.00 30.5	
	ATOM	420	0		A 223	1.306	29.039	21.869	1.00 30.6	
35	ATOM	421	СВ	GLN	A 223	2.434	31.550	20.151	1.00 34.7	
	ATOM	422	CG	GLN	A 223	2.066	32.296	18.873	1.00 38.6	5 C
	ATOM	423	CD	GLN	A 223	3.275	32.719	18.065	1.00 42.4	6 C
	ATOM	424	OE1	GLN	A 223	3.154	33.114	16.903	1.00 45.4	4 0
	MOTA	425	NE2		A 223	4.450	32.652	18.679	1.00 44.9	7 N
40	MOTA	426	N	LEU	A 224	2.226	30.610	23.195	1.00 28.6	4 N
	ATOM	427	CA		A 224	2.632	29.654	24.232	1.00 27.0	
	MOTA	428	С		A 224	3.209	28.401	23.569	1.00 26.4	
	ATOM	429	0		A 224	2.898	27.274	23.962	1.00 25.8	
. –	ATOM	430	CB		A 224	1.424	29.276	25.102	1.00 27.7	
45	MOTA	431	CG		A 224	0.785	30.424	25.893	1.00 27.8	
	MOTA	432			A 224	-0.463	29.931	26.615	1.00 29.5	
	MOTA	433			A 224	1.789	30.981	26.884	1.00 27.5	
	ATOM	434	N		A 225	4.071	28.614	22.577	1.00 25.7	
50	ATOM	435	CA		A 225	4.667	27.531	21.798	1.00 25.8	
50	ATOM	436	C		A 225	5.454	26.473	22.563	1.00 25.3	
	MOTA	437	O CB		A 225 A 225	5.446 5.557	25.302	22.182	1.00 25.0	
	MOTA	438					28.110	20.696	1.00 26.3	
	ATOM	439 440	OG N		A 225 A 226	6.710 6.132	28.731 26.880	21.233	1.00 29.3	
55	ATOM	441	CA		A 226	6.132	25.948	23.630	1.00 24.5	
33	ATOM	442	CA		A 226		25.348	24.424	1.00 24.	
	ATOM ATOM	443	0		A 226		24.529	25.631 26.336	1.00 24.0	
	ATOM	444	СВ		A 226		26.629	24.905	1.00 24.	
	WIOM	-2-73-2	CD	LIE I	A 440	0.219	40.043	44.700	1.00 24.	70 C

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	ATOM	445	CG	MET A 226	9.329	26.715	23.870		C
	MOTA	446		MET A 226	9.960	25.094	23.351	1.00 27.27	S
	ATOM	447	CE	MET A 226	10.773	24.531	24.858 25.872		C
5	ATOM	448 449	N CA	LEU A 227 LEU A 227	4.969 4.225	25.850 25.377	27.030		N C
J	ATOM ATOM	450	C	LEU A 227	3.882	23.377	27.030	1.00 23.65	C
	ATOM	451	Ö	LEU A 227	4.062	23.218	28.052	1.00 24.44	ō
	MOTA	452	СВ	LEU A 227	2.949	26.212	27.237	1.00 24.02	c
	ATOM	453	CG	LEU A 227	2.139	25.868	28.494	1.00 24.67	Č
10	ATOM	454		LEU A 227	3.019	25.994	29.730	1.00 25.75	C
	ATOM	455	-	LEU A 227	0.936	26.798	28.612	1.00 25.81	C
	ATOM	456	N	PRO A 228	3.395	23.336	25.901	1.00 24.00	N
	ATOM	457	CA	PRO A 228	3.073	21.904	25.931	1.00 23.78	С
	MOTA	458	С	PRO A 228	4.261	21.024	26.330	1.00 23.69	С
15	MOTA	459	0	PRO A 228	4.123	20.109	27.155	1.00 23.20	0
	MOTA	460	CB	PRO A 228	2.602	21.626	24.504	1.00 24.23	С
	MOTA	461	CG	PRO A 228	1.957	22.939	24.110	1.00 24.58	С
	MOTA	462	CD	PRO A 228	2.962	23.948	24.629	1.00 23.63	С
	MOTA	463	N	HIS A 229	5.421	21.305	25.747	1.00 22.38	N
20	MOTA	464	CA	HIS A 229	6.626	20.532	26.037	1.00 22.16	С
	MOTA	465	С	HIS A 229	7.089	20.679	27.490	1.00 21.32	С
	MOTA	466	0	HIS A 229	7.409	19.687	28.151	1.00 20.38	0
	MOTA	467	CB	HIS A 229	7.765	20.951	25.103	1.00 22.65	C
05	ATOM	468	CG	HIS A 229	9.037	20.196	25.337	1.00 23.54	C
25	MOTA	469		HIS A 229	9.235	18.910	24.883	1.00 24.88	N
	MOTA	470		HIS A 229	10.160	20.535	26.012	1.00 23.99 1.00 25.42	C C
	ATOM	471		HIS A 229 HIS A 229	10.427	18.488 19.455	25.270 25.957	1.00 23.42	N
	MOTA MOTA	472 473	Nez N	LEU A 230	11.009 7.139	21.913	27.985	1.00 23.32	N
30	MOTA	474	CA	LEU A 230	7.578	22.139	29.355	1.00 20.30	C
30	ATOM	475	C	LEU A 230	6.563	21.623	30.361	1.00 21.22	C
	ATOM	476	Ö	LEU A 230	6.938	21.164	31.435	1.00 19.50	ŏ
	ATOM	477	СВ	LEU A 230	7.858	23.625	29.602	1.00 21.98	Ċ
	ATOM	478	CG	LEU A 230	9.051	24.211	28.839	1.00 23.32	C
35	ATOM	479	CD1	LEU A 230	9.285	25.637	29.322	1.00 25.98	С
	ATOM	480	CD2	LEU A 230	10.311	23.371	29.073	1.00 24.89	С
	MOTA	481	N	ALA A 231	5.279	21.703	30.022	1.00 20.40	N
	MOTA	482	CA	ALA A 231	4.243	21.197	30.917	1.00 21.50	С
	MOTA	483	С	ALA A 231	4.421	19.685	31.040	1.00 21.12	С
40	MOTA	484	0	ALA A 231	4.303	19.124	32.129	1.00 21.78	0
	MOTA	485	CB	ALA A 231	2.859	21.522	30.361	1.00 22.95	C
	MOTA	486	N	ASP A 232	4.707	19.028	29.919	1.00 20.87	N
	MOTA	487	CA	ASP A 232	4.910	17.582	29.916	1.00 21.48	C
45	ATOM	488	C	ASP A 232	6.168	17.228	30.711	1.00 20.43	C
45	MOTA	489	0	ASP A 232	6.167	16.259	31.463	1.00 21.59	0
	MOTA	490	CB	ASP A 232	5.022	17.056	28.482	1.00 21.87	С
	MOTA	491	CG	ASP A 232	3.664	16.893	27.807	1.00 25.14	С
	MOTA	492		ASP A 232	3.639	16.665	26.582	1.00 26.65	0
ΕO	ATOM	493		2 ASP A 232	2.623	16.982	28.497	1.00 25.49	0
50	ATOM	494		LEU A 233	7.228	18.018 17.785	30.549 31.278	1.00 21.20 1.00 20.50	N
	MOTA	495 496		LEU A 233 LEU A 233	8.483 8.267	17.765	32.785	1.00 20.58	C
	MOTA			LEU A 233		17.139	33.587	1.00 20.38	
	MOTA MOTA	497 498		LEU A 233		18.770	30.811	1.00 18.39	C
55	MOTA	499		LEU A 233			31.684	1.00 20.92	C
JJ	ATOM	500		LEU A 233			31.652		C
	ATOM	501		2 LEU A 233					C
	ATOM	502		VAL A 234					Ŋ
	RION		••	*******************		10.501		2.00 20.07	-7

	MOTA	503		VAL A 234	7.263	19.217	34.583	1.00 20.15	С
	ATOM	504		VAL A 234	6.320	18.152	35.146	1.00 19.97	C
	MOTA	505	-	VAL A 234	6.500	17.691	36.268	1.00 19.99	0
_	MOTA	506		VAL A 234	6.665	20.630	34.796	1.00 21.02	C
5	MOTA	507		VAL A 234	6.104	20.778	36.209	1.00 23.20	C
	MOTA	508		VAL A 234	7.754	21.679	34.566	1.00 21.83	C
	MOTA	509	N	SER A 235	5.324	17.749	34.362	1.00 18.23	N
	MOTA	510	CA	SER A 235	4.378	16.732	34.821	1.00 19.68	С
40	MOTA	511	C	SER A 235	5.117	15.413	35.079	1.00 19.46	C
10	MOTA	512	0	SER A 235	4.906	14.743	36.095	1.00 19.95	0
	MOTA	513	CB	SER A 235	3.284	16.537	33.767	1.00 21.15	C
	ATOM	514	OG	SER A 235 TYR A 236	2.229	15.734	34.274	1.00 26.38	0
	ATOM	515 516	N	TYR A 236	5.983	15.057	34.140	1.00 19.05 1.00 19.13	N C
15	MOTA	516 517	CA		6.796	13.849	34.222		
15	ATOM	517 518	C	TYR A 236 TYR A 236	7.660 7.792	13.930 12.958	35.479 36.223	1.00 18.86 1.00 18.39	С О
	MOTA	519	O CB	TYR A 236	7.675	13.781	32.976	1.00 18.39	C
	MOTA MOTA	520	CG	TYR A 236	8.800	12.764	32.970	1.00 19.07	C
	ATOM	521	CD1		8.601	11.466	32.527	1.00 19.18	C
20	ATOM	522	CD2		10.084	13.131	33.391	1.00 20.55	C
20	ATOM	523	CE1		9.665	10.557	32.448	1.00 20.33	C
	ATOM	524	CE2		11.149	12.233	33.321	1.00 21.40	C
	ATOM	525	CZ	TYR A 236	10.934	10.954	32.846	1.00 20.00	c
	ATOM	526	OH	TYR A 236	11.996	10.079	32.749	1.00 21.78	Ö
25	ATOM	527	N	SER A 237	8.241	15.105	35.711	1.00 17.93	N
20	ATOM	528	CA	SER A 237	9.106	15.312	36.868	1.00 18.19	c
	MOTA	529	C	SER A 237	8.373	15.218	38.199	1.00 18.73	Č
	ATOM	530	ō	SER A 237	8.929	14.737	39.184	1.00 19.34	ō
	MOTA	531	СВ	SER A 237	9.830	16.654	36.730	1.00 18.72	Č
30	ATOM	532	OG	SER A 237	10.648	16.628	35.573	1.00 19.76	0
	ATOM	533	N	ILE A 238	7.128	15.680	38.237	1.00 18.89	N
	ATOM	534	CA	ILE A 238	6.343	15.597	39.460	1.00 20.25	С
	ATOM	535	С	ILE A 238	6.101	14.119	39.759	1.00 20.17	С
	ATOM	536	0	ILE A 238	6.129	13.705	40.914	1.00 20.62	0
35	MOTA	537	CB	ILE A 238	4.984	16.337	39.317	1.00 21.21	С
	MOTA	538	CG1	ILE A 238	5.226	17.847	39.236	1.00 23.61	С
	MOTA	539	CG2	ILE A 238	4.068	16.001	40.502	1.00 23.76	С
	MOTA	540	CD1	ILE A 238	3.972	18.668	38.937	1.00 24.70	С
	MOTA	541	N	GLN A 239	5.868	13.315	38.719	1.00 20.04	N
40	MOTA	542	CA	GLN A 239	5.657	11.890	38.936	1.00 19.72	С
	MOTA	543	С	GLN A 239	6.911	11.261	39.531	1.00 20.24	С
	MOTA	544	0	GLN A 239	6.823				0
	MOTA	545	СВ	GLN A 239	5.288	11.178	37.628	1.00 21.35	С
	ATOM	546	CG	GLN A 239	3.920	11.576	37.086	1.00 21.87	С
45	MOTA	547	CD	GLN A 239	3.487	10.707	35.922	1.00 23.58	С
	ATOM	548		GLN A 239	3.092	9.556	36.105	1.00 26.39	0
	MOTA	549		GLN A 239	3.568	11.249	34.720	1.00 22.31	N
	MOTA	550	N	LYS A 240	8.080	11.661	39.037	1.00 19.37	N
	MOTA	551	CA	LYS A 240	9.336	11.116	39.557	1.00 19.49	C
50	ATOM	552	С	LYS A 240	9.575	11.583	40.994	1.00 20.03	C
	MOTA	553	0	LYS A 240	10.086	10.826	41.826	1.00 20.81	0
	MOTA	554	CB	LYS A 240	10.509	11.525	38.658	1.00 19.27	C
	ATOM	555	CG	LYS A 240	10.385	11.015	37.216	1.00 19.70	C
55	ATOM	556 557	CD	LYS A 240	10.174 10.201	9.491	37.165	1.00 20.85	C
JJ	ATOM	557 558	CE NZ	LYS A 240	9.919	8.986	35.734	1.00 20.78	C
	ATOM	559	NZ N	LYS A 240 VAL A 241	9.919	7.527	35.631	1.00 21.79	N
	ATOM	560	CA	VAL A 241 VAL A 241	9.355	12.827	41.284 42.630		N
	MOTA	200	CA	AWD W 741	9.333	13.380	42.030	1.00 21.18	С

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	MOTA	561	С	VAL A 241	8.466	12.633	43.621	1.00 22.58	С
	ATOM	562	0	VAL A 241	8.845	12.418	44.769	1.00 22.01	0
	MOTA	563	CB	VAL A 241	9.006	14.890	42.658	1.00 22.53	С
	MOTA	564	CG1	VAL A 241	8.893	15.392	44.104	1.00 23.49	С
5	MOTA	565	CG2	VAL A 241	10.092	15.671	41.929	1.00 22.43	С
	MOTA	566	N	ILE A 242	7.277	12.237	43.178	1.00 22.44	N
	ATOM	567	CA	ILE A 242	6.375	11.492	44.052	1.00 23.64	С
	ATOM	568	С	ILE A 242	7.027	10.157	44.416	1.00 23.45	С
	MOTA	569	0	ILE A 242	6.987	9.726	45.573	1.00 25.50	0
10	MOTA	570	CB	ILE A 242	5.012	11.255	43.360	1.00 24.32	С
	MOTA	571	CG1	ILE A 242	4.235	12.575	43.303	1.00 25.64	C
	MOTA	572	CG2	ILE A 242	4.214	10.186	44.104	1.00 24.95	С
	MOTA	573	CD1	ILE A 242	3.012	12.540	42.401	1.00 25.41	С
	ATOM	574	N	GLY A 243	7.652	9.521	43.431	1.00 22.76	N
15	MOTA	575	CA	GLY A 243	8.310	8.246	43.665	1.00 23.14	С
	MOTA	576	С	<b>GLY A 243</b>	9.491	8.385	44.604	1.00 23.29	С
	MOTA	577	0	<b>GLY A 243</b>	9.719	7.525	45.454	1.00 24.26	0
	MOTA	578	N	PHE A 244	10.244	9.471	44.443	1.00 22.21	N
	MOTA	579	CA	PHE A 244	11.406	9.754	45.287	1.00 23.08	C
20	ATOM	580	С	PHE A 244	10.962	9.960	46.734	1.00 23.33	С
	ATOM	581	0	PHE A 244	11.509	9.359	47.665	1.00 22.96	0
	ATOM	582	CB	PHE A 244	12.110	11.023	44.799	1.00 21.55	С
	ATOM	583	CG	PHE A 244	13.264	11.454	45.663	1.00 23.20	С
	MOTA	584	CD1	PHE A 244	14.474	10.764	45.632	1.00 25.04	С
25	ATOM	585	CD2	PHE A 244	13.140	12.548	46.516	1.00 24.78	С
	ATOM	586	CE1	PHE A 244	15.542	11.157	46.437	1.00 25.46	C
	ATOM	587	CE2	PHE A 244	14.205	12.950	47.327	1.00 24.71	С
	MOTA	588	CZ	PHE A 244	15.407	12.254	47.286	1.00 24.22	С
	ATOM	589	N	ALA A 245	9.963	10.819	46.912	1.00 23.25	N
30	ATOM	590	CA	ALA A 245	9.441	11.134	48.233	1.00 23.37	С
	ATOM	591	С	ALA A 245	8.960	9.906	49.006	1.00 25.09	C
	MOTA	592	0	ALA A 245	9.182	9.805	50.212	1.00 24.87	0
	MOTA	593	СВ	ALA A 245	8.310	12.156	48.113	1.00 22.36	С
	ATOM	594	N	LYS A 246	8.309	8.975	48.314	1.00 26.15	N
35	MOTA	595	CA	LYS A 246	7.800	7.768	48.959	1.00 28.66	С
	ATOM	596	С	LYS A 246	8.914	6.918	49.562	1.00 29.21	С
	ATOM	597	0	LYS A 246	8.668	6.117	50.466	1.00 29.75	0
	ATOM	598	CB	LYS A 246	6.997	6.931	47.957	1.00 30.93	С
	MOTA	599	CG	LYS A 246	5.702	7.593	47.501	1.00 34.75	С
40	MOTA	600	CD	LYS A 246	5.017	6.811	46.383	1.00 37.28	С
	ATOM	601	CE	LYS A 246	4.410	5.501	46.873	1.00 40.02	С
	MOTA	602	NZ	LYS A 246	3.230	5.724	47.756	1.00 42.15	N
	ATOM	603	N	MET A 247	10.138	7.104	49.074	1.00 28.68	N
	MOTA	604	CA	MET A 247	11.282	6.339	49.562	1.00 29.45	С
45	MOTA	605	С	MET A 247	12.076	7.021	50.681	1.00 28.75	С
	ATOM	606	0	MET A 247	13.012	6.431	51.230	1.00 28.61	0
	MOTA	607	CB	MET A 247	12.219	5.990	48.396	1.00 30.97	С
	MOTA	608	CG	MET A 247	11.614	5.007	47.393	1.00 34.76	С
	MOTA	609	SD	MET A 247	12.766	4.475	46.096	1.00 39.72	S
50	MOTA	610	CE	MET A 247	12.303	5.554	44.763	1.00 39.07	С
	MOTA	611	N	ILE A 248	11.709	8.253	51.023	1.00 27.31	N
	ATOM	612	CA	ILE A 248	12.391	8.973	52.100	1.00 28.07	С
	ATOM	613	С	ILE A 248	12.033	8.295	53.420	1.00 28.99	С
	ATOM	614	0	ILE A 248	10.859	8.179	53.763	1.00 28.97	0
55	ATOM	615	CB	ILE A 248	11.934		52.195	1.00 27.49	C
	ATOM	616	CG:		12.299		50.916	1.00 26.51	С
	ATOM	617		2 ILE A 248	12.582		53.411	1.00 28.63	C
	MOTA	618		1 ILE A 248	11.730		50.865	1.00 25.47	C

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	MOTA	619	N	PRO A		13.041	7.844	54.181	1.00		N
	MOTA	620	CA	PRO A		12.764	7.182	55.460	1.00		C
	MOTA	621	C	PRO A		11.818	7.992	56.348	1.00		C
_	MOTA	622	0	PRO A		12.107	9.138	56.688	1.00	34.02	0
5	MOTA	623	CB	PRO A		14.153	7.035	56.075	1.00		С
	ATOM	624	CG	PRO A	249	15.021	6.835	54.871	1.00		С
	MOTA	625	CD	PRO A		14.490	7.890	53.917	1.00		С
	MOTA	626	N	GLY A		10.686	7.392	56.706	1.00		N
	MOTA	627	CA	GLY A		9.725	8.064	57.565	1.00		С
10	MOTA	628	С	GLY A		8.542	8.700	56.858	1.00		С
	MOTA	629	0	GLY A		7.484	8.888	57.459		33.45	0
	MOTA	630	N	PHE A		8.709	9.023	55.579		33.51	N
	MOTA	631	CA	PHE A		7.643	9.658	54.809		33.69	С
	MOTA	632	C	PHE A		6.335	8.871	54.833		34.57	C
15	MOTA	633	0	PHE A		5.259	9.455	54.964		35.10	0
	ATOM	634	CB	PHE A		8.082	9.850	53.356		31.35	С
	ATOM	635	CG	PHE A	A 251	7.180	10.754	52.564		29.89	С
	MOTA	636	CD1	PHE A	A 251	7.234	12.134	52.735		30.12	C
	MOTA	637		PHE A		6.276	10.227	51.643		30.05	С
20	MOTA	638	CE1			6.400	12.979	51.999	1.00	29.28	С
	MOTA	639	CE2	PHE A	A 251	5.441	11.063	50.906	1.00	28.78	С
	MOTA	640	CZ	PHE A		5.505	12.440	51.085		28.48	С
	MOTA	641	N	ARG A		6.431	7.551	54.703		36.83	N
	MOTA	642	CA		A 252	5.250	6.691	54.698		39.19	C
25	MOTA	643	С		A 252	4.535	6.647	56.045		39.61	C
	MOTA	644	0		A 252	3.391	6.200	56.127		40.31	0
	MOTA	645	CB		A 252	5.625	5.262	54.292	1.00	41.06	С
	MOTA	646	CG		A 252	6.138	5.101	52.867		44.96	С
	MOTA	647	CD		A 252	6.260	3.620	52.516	1.00	47.63	С
30	MOTA	648	NE	ARG 2	A 252	6.777	3.393	51.169	1.00	50.79	N
	MOTA	649	CZ		A 252	8.062	3.459	50.831		51.79	С
	MOTA	650		ARG		8.982	3.745	51.745		52.82	N
	MOTA	651	NH2		A 252	8.427	3.235	49.576		52.64	N
	MOTA	652	N		A 253	5.205	7.102	57.098		39.96	N
35	MOTA	653	CA		A 253	4.610	7.097	58.430		40.45	С
	MOTA	654	С		A 253	3.648	8.255	58.635	1.00	39.90	С
	MOTA	655	0		A 253	2.902	8.284	59.612	1.00	39.68	0
	MOTA	656	CB		A 253	5.698	7.127	59.506		42.53	С
	MOTA	657	CG		A 253	6.524	5.856	59.531		44.84	С
40	MOTA	658		ASP		5.938	4.767	59.345		47.60	0
	MOTA	659		ASP		7.752	5.942	59.743		45.66	0
	MOTA	660	N		A 254		9.208			38.00	
	MOTA	661	CA		A 254		10.361	57.780		37.81	C
4.5	MOTA	662	С		A 254		9.978	57.218		37.76	С
45	MOTA	663	0		A 254		9.000	56.476		37.49	0
	MOTA	664	CB		A 254		11.521	56.955		36.51	С
	MOTA	665	CG		A 254		12.101	57.346		36.86	С
	MOTA	666		LEU			13.113	56.297		35.66	C
	MOTA	667		LEU			12.751	58.719		36.85	С
50	MOTA	668	Ŋ		A 255		10.745	57.579		38.14	
	MOTA	669	CA		A 255		10.496	57.077		39.08	
	MOTA	670	C		A 255		10.828	55.588		39.76	C
	ATOM	671	0		A 255		11.593	55.139		39.15	
	ATOM	672	CB		A 255		11.397	57.781		39.68	
55	ATOM	673		L THR			12.767	57.677		39.47	
	ATOM	674		2 THR			11.020			40.24	
	MOTA	675	N		A 256		10.258			40.24	
	ATOM	676	CA	SER	A 256	-1.883	10.527	53.382	1.00	40.92	С

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	ATOM	677	С	SER A	A :	256	-2.148	12.010	53.152	1.00		C
	ATOM	678	0	SER A	A	256	-1.662	12.599	52.185	1.00	40.28	0
	ATOM	679	CB	SER A	A	256	-2.968	9.693	52.690	1.00	41.53	С
	ATOM	680	OG	SER .	A	256	-4.263	10.203	52.957	1.00	43.52	0
5	ATOM	681	N	GLU 2	A	257	-2.916	12.610	54.056	1.00	39.55	N
	ATOM	682	CA	GLU .	A	257	-3.252	14.024	53.963	1.00	38.71	С
	ATOM	683	С	GLU .	A	257	-1.999	14.889	54.038	1.00	36.50	C
	MOTA	684	0	GLU .	Α	257	-1.825	15.810	53.240	1.00	36.20	0
	MOTA	685	СВ	GLU			-4.221	14.400	55.085		41.43	C
10	ATOM	686	CG	GLU			-4.650	15.853	55.090		44.63	C
	ATOM	687	CD	GLU			-5.747	16.121	56.103		47.39	C
	ATOM	688		GLU			-6.879	15.634	55.896		48.76	ō
	ATOM	689	OE2	GLU			-5.476	16.810	57.109		48.99	ŏ
	ATOM	690	N	ASP			-1.132	14.593	55.001		34.59	N
15	ATOM	691	CA	ASP			0.111	15.339	55.159		33.04	C
13			CA	ASP								
	MOTA	692					1.064	15.047	54.002		32.48	C
	ATOM	693	0	ASP			1.782	15.934	53.546		31.37	0
	МОТА	694	CB	ASP			0.784	14.984	56.488		34.07	C
00	MOTA	695	CG	ASP			0.256	15.809	57.645		35.11	C
20	MOTA	696		ASP			0.599	15.501	58.807		35.63	0
	MOTA	697		ASP			-0.493	16.775	57.386		34.86	0
	MOTA	698	N	GLN			1.072	13.803	53.532		31.90	N
	MOTA	699	CA	GLN			1.940	13.433	52.417	1.00	32.81	С
	MOTA	700	С	GLN			1.611	14.272	51.184	1.00	32.59	С
25	MOTA	701	0	GLN	A	259	2.505	14.820	50.534	1.00	32.51	0
	MOTA	702	CB	GLN	Α	259	1.783	11.946	52.077		32.98	С
	ATOM	703	CG	GLN	Α	259	2.217	11.000	53.181	1.00	34.94	С
	MOTA	704	CD	GLN	Α	259	2.168	9.547	52.755	1.00	37.19	С
	ATOM	705	OE1	GLN	Α	259	2.322	8.641	53.576	1.00	39.55	0
30	ATOM	706	NE2	GLN	Α	259	1.958	9.315	51.466	1.00	37.81	N
	MOTA	707	N	ILE	Α	260	0.325	14.375	50.866		32.68	N
	MOTA	708	CA	ILE	Α	260	-0.109	15.147	49.706			С
	ATOM	709	С			260	0.183	16.634	49.880	1.00		C
	ATOM	710	0			260	0.588	17.311	48.933			Ō
35	ATOM	711	СВ			260	-1.619	14.959	49.445		33.97	C
•	ATOM	712	CG1			260	-1.933	13.471	49.277	1.00	34.59	C
	ATOM	713	CG2			260	-2.036	15.731	48.201	1.00		C
	ATOM	714	CD1			260	-1.156	12.789	48.165	1.00		Ċ
	ATOM	715	N			261	-0.029	17.146	51.088		29.87	N
40	ATOM	716	CA			261	0.244	18.551	51.358		28.91	C
70	ATOM	717	C			261	1.717	18.862	51.097		27.91	C
	ATOM	718	Ö			261	2.043	19.856	50.445		28.13	o
		719				261	-0.089	18.923	52.827		28.91	
	MOTA		CB								30.09	C
45	ATOM	720		VAL			0.472	20.294	53.161			C
45	ATOM	721		VAL			-1.594	18.911	53.035		31.46	C
	MOTA	722	N			262	2.605	18.011	51.604		27.17	N
	MOTA	723	CA			262	4.039	18.222	51.423		25.81	C
	ATOM	724	C			262	4.461	18.126	49.955		25.46	C
	ATOM	725	0			262	5.274	18.921	49.485		24.78	0
50	MOTA	726	СВ			262	4.836	17.219	52.265		26.02	C
	MOTA	727	CG			262	4.604	17.278	53.781		25.71	С
	MOTA	728		. LEU			5.382	16.162	54.464		26.98	С
	MOTA	729		LEU			5.028	18.634	54.317		26.30	С
	MOTA	730	N			263	3.911	17.155	49.232	1.00	25.71	N
55	MOTA	731	CA	LEU	Α	263	4.244	16.985	47.818	1.00	26.15	С
	ATOM	732	С	LEU	Α	263	3.763	18.166	46.974	1.00	26.42	С
	ATOM	733	0	LEU	Α	263	4.514	18.702	46.154	1.00	25.91	0
	ATOM	734	СВ	LEU	Α	263	3.633	15.681	47.283		27.20	С

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	ATOM	735		LEU A 263	4.293	14.376	47.745	1.00 29.34	С
	MOTA	736	CD1	LEU A 263	3.401	13.197	47.404	1.00 30.14	С
	MOTA	737	CD2	LEU A 263	5.658	14.223	47.082	1.00 31.00	С
	ATOM	738	N	LYS A 264	2.519	18.585	47.178	1.00 25.69	N
5	MOTA	739	CA	LYS A 264	1.987	19.699	46.405	1.00 26.59	C
	MOTA	740	С	LYS A 264	2.709	21.011	46.655	1.00 26.97	С
	MOTA	741	0	LYS A 264	2.962	21.767	45.723	1.00 27.99	0
	ATOM	742	CB	LYS A 264	0.496	19.899	46.688	1.00 29.36	C
	MOTA	743	CG	LYS A 264	-0.417	18.910	45.994	1.00 31.84	C
10	ATOM	744	CD	LYS A 264	-1.862	19.348	46.156	1.00 35.05	С
	ATOM	745	CE	LYS A 264	-2.822	18.400	45.468	1.00 37.95	C
	MOTA	746	NZ	LYS A 264	-4.233	18.872	45.629	1.00 39.41	N
	ATOM	747	N	SER A 265	3.047	21.286	47.908	1.00 26.64	N
	ATOM	748	CA	SER A 265	3.712	22.540	48.227	1.00 27.75	C
15	ATOM	749	С	SER A 265	5.199	22.591	47.884	1.00 26.92	С
	ATOM	750	0	SER A 265	5.750	23.676	47.723	1.00 28.28	0
	MOTA	751	CB	SER A 265	3.513	22.881	49.709	1.00 28.81	С
	MOTA	752	OG	SER A 265	4.101	21.902	50.540	1.00 33.64	0
	MOTA	753	N	SER A 266	5.847	21.434	47.757	1.00 25.17	N
20	MOTA	754	CA	SER A 266	7.277	21.412	47.449	1.00 23.91	С
	MOTA	755	С	SER A 266	7.609	21.035	46.011	1.00 23.51	C
	MOTA	756	0	SER A 266	8.749	21.206	45.572	1.00 23.30	0
	ATOM	757	CB	SER A 266	8.001	20.445	48.385	1.00 24.45	C
	ATOM	758	OG	SER A 266	7.656	19.101	48.094	1.00 24.60	0
25	ATOM	759	N	ALA A 267	6.619	20.519	45.285	1.00 22.67	N
	MOTA	760	CA	ALA A 267	6.801	20.089	43.898	1.00 23.39	С
	MOTA	761	С	ALA A 267	7.698	20.979	43.040	1.00 23.51	С
	ATOM	762	0	ALA A 267	8.716	20.517	42.515	1.00 23.55	0
	MOTA	763	CB	ALA A 267	5.436	19.938	43.217	1.00 24.51	C
30	MOTA	764	N	ILE A 268	7.330	22.247	42.883	1.00 22.01	N
	ATOM	765	CA	ILE A 268	8.132	23.135	42.041	1.00 22.23	С
	MOTA	766	С	ILE A 268	9.539	23.374	42.592	1.00 22.05	С
	MOTA	767	0	ILE A 268	10.494	23.558	41.828	1.00 20.90	0
	MOTA	768	CB	ILE A 268	7.426	24.496	41.811	1.00 23.63	С
35	MOTA	769	CG1	ILE A 268	8.097	25.232	40.645	1.00 24.85	С
	MOTA	770	CG2	ILE A 268	7.484	25.354	43.068	1.00 25.18	С
	ATOM	771	CD1		7.933	24.549	39.303	1.00 25.69	С
	MOTA	772	N	GLU A 269	9.674	23.352	43.911	1.00 20.40	N
	MOTA	773	CA	GLU A 269	10.979	23.561	44.529	1.00 20.63	С
40	MOTA	774	С	GLU A 269	11.933	22.402	44.268	1.00 21.33	С
	MOTA	775	0	GLU A 269	13.109	22.620	43.976	1.00 20.99	0
	MOTA	776	CB	GLU A 269	10.823	23.770	46.030		С
	MOTA	777	CG	GLU A 269	10.206	25.110	46.396	1.00 22.10	С
	MOTA	778	CD	GLU A 269	10.009	25.261	47.892	1.00 23.72	С
45	MOTA	779		GLU A 269	10.803	24.670	48.656	1.00 22.73	0
	ATOM	780	OE2	GLU A 269	9.067	25.974	48.301	1.00 24.58	0
	MOTA	781	N	VAL A 270	11.434	21.172	44.375	1.00 20.39	N
	MOTA	782	CA	VAL A 270	12.279	20.006	44.143	1.00 20.83	С
	MOTA	783	С	VAL A 270	12.644	19.911	42.670	1.00 20.52	С
50	MOTA	784	0	VAL A 270	13.734	19.458	42.318	1.00 20.87	0
	MOTA	785	CB	VAL A 270	11.582	18.709	44.597	1.00 21.55	
	ATOM	786		VAL A 270	12.481	17.512	44.318	1.00 21.95	
	ATOM	787		VAL A 270	11.268	18.790	46.086	1.00 23.25	С
	MOTA	788	N	ILE A 271	11.731	20.337	41.804	1.00 20.29	N
55	MOTA	789	CA	ILE A 271	12.010	20.318	40.376	1.00 20.71	
	MOTA	790	С	ILE A 271	13.145	21.300	40.099	1.00 20.86	
	MOTA	791	0	ILE A 271	14.083	20.990	39.361	1.00 20.78	
	ATOM	792	CB	ILE A 271	10.755	20.684	39.563	1.00 21.89	С

	MOTA	793	CG1	ILE A			9.842	19.450	39.483	1.00		С
	MOTA	794	CG2	ILE A	A	271	11.149	21.173	38.170	1.00		С
	MOTA	795	CD1	ILE A	Α	271	8.489	19.711	38.852	1.00	27.85	С
	ATOM	796	N	MET .	A	272	13.076	22.481	40.701	1.00	21.17	N
5	MOTA	797	CA	MET .	A	272	14.147	23.446	40.500	1.00	21.57	С
	ATOM	798	С	MET .	A	272	15.474	22.888	41.020	1.00	20.82	С
	MOTA	799	0	MET .	Α	272	16.513	23.064	40.384	1.00	22.20	0
	ATOM	800	CB	MET .	Α	272	13.800	24.770	41.183		22.31	С
	ATOM	801	CG	MET			12.595	25.441	40.549		24.16	C
10	ATOM	802	SD	MET			12.222	27.036	41.296		26.22	s
. •	ATOM	803	CE	MET			11.003	27.687	40.134		26.38	č
	ATOM	804	N	LEU			15.442	22.204	42.163		21.17	N
	ATOM	805	CA	LEU			16.661	21.606	42.717		21.28	Ĉ
	ATOM	806	C	LEU			17.226	20.486	41.842		20.96	c
15	ATOM	807	Ö	LEU			18.408	20.494	41.487		20.75	o
13								21.026	44.116		22.98	C
	MOTA	808	CB	LEU			16.405					
	ATOM	809	CG	LEU			16.367	21.940	45.337		25.62	C
	ATOM	810	CD1	LEU			15.959	21.129	46.572		25.83	C
~~	ATOM	811	CD2	LEU			17.736	22.571	45.543		26.65	С
20	ATOM	812	N	ARG			16.385	19.517	41.494		19.69	N
	MOTA	813	CA	ARG			16.852	18.384	40.702		19.52	С
	MOTA	814	С	ARG			17.317	18.787	39.309		19.10	С
	MOTA	815	0	ARG			18.159	18.117	38.715		19.83	0
	ATOM	816	CB	ARG			15.759	17.299	40.610		19.75	С
25	MOTA	817	CG	ARG			14.652	17.566	39.601		19.52	С
	MOTA	818	CD	ARG	Α	274	13.381	16.792	39.969	1.00	19.72	С
	MOTA	819	NE	ARG	A	274	13.599	15.356	40.153	1.00	18.11	N
	MOTA	820	CZ	ARG	Α	274	13.580	14.453	39.175	1.00	19.01	С
	MOTA	821	NH1	ARG	Α	274	13.357	14.824	37.919	1.00	18.53	N
30	MOTA	822	NH2	ARG	Α	274	13.759	13.168	39.458	1.00	19.51	N
	MOTA	823	N			275	16.792	19.892	38.793	1.00	19.73	N
	ATOM	824	CA			275	17.183	20.331	37.463	1.00		С
	ATOM	825	С			275	18.615	20.838	37.442		19.90	С
	ATOM	826	0			275	19.191	21.016	36.377		20.21	0
35	ATOM	827	СВ			275	16.249	21.437	36.958		20.51	Č
-	ATOM	828	OG			275	16.520	22.680	37.579		20.38	ō
	ATOM	829	N			276	19.198	21.055	38.615		20.28	N
	ATOM	830	CA			276	20.564	21.557	38.662		19.85	C
	ATOM	831	C			276	21.512	20.544	38.024		21.26	č
40	ATOM	832	Ö			276	22.585	20.903	37.538	1.00		ŏ
70	ATOM	833	СВ			276	20.983	21.843	40.108		20.77	c
	ATOM	834	CG			276		22.651	40.187		23.39	C
	MOTA	835					23.275	22.187	40.713		26.18	-
		836		ASN			22.231	23.867	39.649		21.92	O N
15	MOTA			ASN								N
45	ATOM	837	N			277	21.096	19.280	38.000		20.52	N
	ATOM	838	CA			277	21.925	18.226	37.425		21.75	C
	ATOM	839	С			277	22.103	18.370	35.908		21.79	C
	ATOM	840	0			277	23.105	17.910	35.351		22.41	0
50	ATOM	841	CB			277	21.331	16.852	37.785		22.91	С
50	MOTA	842	CG			277	22.199	15.659	37.413		26.24	C
	MOTA	843	CD			277	21.904	14.418	38.261		28.07	С
	MOTA	844	OE:			277	22.359	13.319	37.875		30.43	0
	MOTA	845		2 GLU			21.233	14.532	39.317		26.56	0
	MOTA	846	N			278	21.152	19.011	35.234		19.68	N
55	ATOM	847	CA			278			33.789	1.00	20.64	С
	MOTA	848	С			278			33.448	1.00	21.58	С
	ATOM	849	0	SER	P	278	22.008	20.910	32.292	1.00	22.05	
	MOTA	850	CB	SER	A	278	19.934	18.910	33.092		20.93	C

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	MOTA	851	OG	SER A			18.941	19.829	33.497		22.00	0
	MOTA	852	N	PHE A			21.751	21.474	34.451	1.00	21.92	N
	MOTA	853	CA	PHE A			22.160	22.853	34.219		23.24	C
_	MOTA	854	С	PHE A			23.659	22.912	33.972	1.00	24.55	С
5	MOTA	855	0	PHE A			24.429	22.218	34.638	1.00	24.49	0
	MOTA	856	CB	PHE A	Ą	279	21.820	23.723	35.429	1.00	23.08	С
	MOTA	857	CG	PHE A			22.051	25.187	35.198	1.00	24.02	С
	MOTA	858	CD1				21.135	25.942	34.471	1.00	24.96	С
	MOTA	859	CD2	PHE A			23.197	25.805	35.682	1.00	24.94	С
10	MOTA	860	CE1	PHE A	A	279	21.356	27.293	34.227	1.00	24.93	С
	MOTA	861	CE2	PHE A			23.429	27.160	35.442		25.50	C
	MOTA	862	CZ	PHE A			22.506	27.903	34.714		24.47	С
	MOTA	863	N	THR A			24.077	23.728	33.010	1.00	24.73	N
	ATOM	864	CA	THR .	A	280	25.496	23.872	32.728	1.00	26.87	C
15	MOTA	865	С	THR :			25.884	25.343	32.672		27.44	С
	MOTA	866	0	THR .			25.186	26.162	32.070		26.28	0
	MOTA	867	CB	THR .			25.897	23.198	31.399	1.00	27.76	С
	ATOM	868	OG1				27.298	23.408	31.173	1.00	31.72	0
	ATOM	869	CG2	THR	A	280	25.107	23.768	30.236		27.79	С
20	ATOM	870	N	MET			26.991	25.676	33.326	1.00	28.33	N
	ATOM	871	CA	MET			27.469	27.049	33.340	1.00	31.03	С
	MOTA	872	С	MET	A	281	28.275	27.390	32.095	1.00	31.28	С
	ATOM	873	0	MET	A	281	28.812	28.490	31.980	1.00	30.87	0
	MOTA	874	CB	MET			28.298	27.306	34.596	1.00	33.43	C
25	MOTA	875	CG	MET	A	281	27.448	27.518	35.835	1.00	36.11	С
	MOTA	876	SD	MET			28.429	27.829	37.295	1.00	39.85	S
	MOTA	877	CE	MET			28.995	29.495	36.967	1.00	40.40	С
	MOTA	878	N	ASP	Α	282	28.364	26.448	31.159	1.00	31.72	N
	MOTA	879	CA	ASP	Α	282	29.097	26.709	29.925	1.00	32.91	С
30	MOTA	880	С	ASP			28.366	27.818	29.175	1.00	32.02	С
	MOTA	881	0	ASP	Α	282	28.989	28.764	28.683	1.00	31.15	0
	MOTA	882	CB	ASP	Α	282	29.172	25.455	29.050	1.00	35.93	C
	ATOM	883	CG	ASP			29.947	24.328	29.708		39.91	С
22	MOTA	884	OD1				30.940	24.619	30.412		42.35	0
35	ATOM	885	OD2				29.573	23.150	29.508	1.00	42.45	0
	MOTA	886	N	ASP	Α	283	27.041	27.702	29.100	1.00	29.87	N
	MOTA	887	CA	ASP			26.224	28.704	28.418		28.59	С
	MOTA	888	С	ASP			24.931	29.032	29.170		27.92	С
	MOTA	889	0	ASP			23.984	29.568	28.592		27.21	0
40	MOTA	890	CB	ASP			25.904	28.243	26.994		29.84	С
	ATOM .	891	CG			283	25.030	27.006	26.958		31.11	С
	MOTA	892		. ASP			24.872		28.009		28.99	0
	MOTA	893		ASP			24.507	26.687	25.870		32.79	0
	MOTA	894	N			284	24.902	28.708	30.460		26.84	N
45	ATOM	895	CA			284	23.748	28.985	31.317		27.62	С
	MOTA	896	С			284	22.449	28.379	30.801		27.20	C
	MOTA	897	0			284	21.429	29.060	30.686		27.86	0
	MOTA	898	СВ			284	23.565	30.497	31.484		29.95	С
	MOTA	899	CG			284		31.219	32.031		33.34	С
50	MOTA	900	SD			284		30.578	33.624		36.23	S
	MOTA	901	CE			284		31.242	33.719		35.78	C
	MOTA	902	N			285		27.091	30.503		25.58	N
	MOTA	903	CA			285		26.427	30.010		24.62	
	MOTA	904	C			285		25.090	30.697		24.72	
55	MOTA	905	0			285		24.641	31.415		24.18	
	MOTA	906	CB			. 285		26.186			24.98	
	MOTA	907	OG			285		25.224			25.94	
	MOTA	908	N	TRP	A	286	19.982	24.472	30.480	1.0	24.17	N

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	ATOM	909 910	CA C	TRP A			19.699 19.842	23.146 22.312	30.997 29.732		24.74	C
	MOTA MOTA	911	0	TRP A			19.842	22.312	28.828		25.34 25.37	С
	ATOM	912	СВ	TRP A			18.268	23.064	31.522		23.76	O C
5	ATOM	913	CG	TRP 2			18.048	23.702	32.863		21.76	C
•	ATOM	914		TRP A			18.186	23.107	34.088		21.47	C
	ATOM	915	CD2				17.568	25.031	33.118		23.03	Č
	ATOM	916	NE1	TRP A			17.811	23.976	35.084		21.88	N
	ATOM	917	CE2	TRP			17.429	25.164	34.519		22.96	C
10	ATOM	918	CE3	TRP 2			17.238	26.121	32.299		23.54	Č
	ATOM	919	CZ2	TRP 2			16.970	26.341	35.120		24.15	Ċ
	MOTA	920	CZ3	TRP 2	A 2	86	16.781	27.293	32.898		22.92	С
	ATOM	921	CH2	TRP :	A 2	86	16.651	27.390	34.297	1.00	23.66	С
	ATOM	922	N	THR :	A 2	87	20.918	21.540	29.654	1.00	25.53	N
15	MOTA	923	CA	THR .	A 2	87	21.173	20.721	28.478	1.00	27.14	С
	ATOM	924	С	THR .	A 2	87	20.833	19.266	28.753	1.00	27.53	С
	MOTA	925	0	THR .	A 2	87	21.501	18.607	29.551	1.00	27.40	0
	ATOM	926	CB	THR	A 2	87	22.644	20.853	28.049	1.00	27.77	С
	MOTA	927	OG1	THR			22.914	22.229	27.733		30.32	0
20	MOTA	928	CG2	THR			22.922	20.000	26.824		29.69	С
	MOTA	929	N	CYS			19.792	18.775	28.084		28.08	N
	MOTA	930	CA	CYS			19.326	17.406	28.270		30.34	С
	MOTA	931	C	CYS			19.478	16.520	27.040		33.66	С
05	ATOM	932	0	CYS			18.530	15.857	26.624		33.19	0
25	MOTA	933	CB	CYS			17.861	17.426	28.699		29.32	C
	MOTA	934	SG	CYS			17.566	18.403	30.188		28.01	S
	MOTA	935	N	GLY			20.675	16.498	26.466		37.69	N
	ATOM	936	CA	GLY			20.897	15.682	25.286		41.85	C
30	MOTA	937	С	GLY			21.072	16.536	24.044		44.11	C
30	MOTA	938 939	0	GLY			21.842 20.349	17.497 16.205	24.051		45.10	0
	MOTA	940	N CA	ASN ASN			20.349	16.205	22.978 21.737		46.33 47.32	N
	MOTA MOTA	941	C	ASN			19.961	18.391	21.737		47.22	C
	ATOM	942	Ö	ASN			19.303	18.746	22.857		47.49	0
35	ATOM	943	СВ	ASN			19.733	16.241	20.600		49.56	C
00	ATOM	944	CG	ASN			18.235	16.224	20.792		51.07	C
	ATOM	945	OD1				17.591	17.271	20.803		52.29	ō
	ATOM	946	ND2				17.668	15.032	20.944		51.45	N
	ATOM	947	N	GLN			20.277	19.205	20.874		46.24	N
40	MOTA	948	CA	GLN			19.896	20.611	20.850		45.60	C
	ATOM	949	С	GLN			18.402	20.859	21.031		43.20	C
	MOTA	950	0	GLN			18.007	21.916	21.520	1.00	43.27	0
	ATOM	951	СВ	GLN			20.380	21.247	19.545		47.46	С
	ATOM	952	CG	GLN			21.879	21.087	19.325		50.94	С
45	ATOM	953	CD	GLN	A 2	291	22.705	21.786	20.395	1.00	52.59	C
	MOTA	954	OE1	GLN	A :	291	23.893	21.503	20.563	1.00	54.12	0
	ATOM	955	NE2	GLN	Α :	291	22.081	22.712	21.114	1.00	53.69	N
	MOTA	956	N	ASP	A :	292	17.574	19.897	20.636	1.00	40.92	N
	MOTA	957	CA	ASP			16.129	20.046	20.780		38.58	С
50	MOTA	958	С	ASP			15.740	20.140	22.252		35.80	С
	ATOM	959	0	ASP			14.769	20.814	22.601		34.04	
	MOTA	960	CB	ASP			15.391	18.862	20.145		41.69	С
	MOTA	961	CG	ASP			15.325	18.950	18.629		44.13	С
ce	ATOM	962		ASP			14.862	17.973	18.002		45.48	
55	MOTA	963		ASP			15.724	19.993	18.067		45.78	
	MOTA	964	N	TYR			16.506	19.469	23.111		33.09	
	ATOM	965	CA	TYR			16.219	19.465	24.543		31.43	
	MOTA	966	С	TYR	A	293	17.183	20.305	25.367	1.00	30.28	С

		0.65	_	m	000	10 550	10 004	06 404	1 00 30 50	_
	ATOM	967	0	TYR A		17.558	19.934	26.481	1.00 30.56	0
	MOTA	968	CB	TYR A		16.186	18.027	25.066	1.00 31.64	С
	MOTA	969	CG	TYR A		15.232	17.154	24.287	1.00 31.43	C
_	MOTA	970		TYR A		15.591	15.864	23.905	1.00 32.22	C
5	MOTA	971		TYR A		13.999	17.647	23.861	1.00 32.09	С
	MOTA	972		TYR A		14.752	15.091	23.106	1.00 32.84	С
	MOTA	973		TYR A		13.153	16.883	23.063	1.00 31.78	С
	MOTA	974	CZ	TYR A		13.537	15.611	22.684	1.00 33.14	С
	MOTA	975	OH	TYR A	293	12.726	14.874	21.850	1.00 32.75	0
10	MOTA	976	N	LYS A	294	17.594	21.431	24.801	1.00 29.44	N
	MOTA	977	CA	LYS A	294	18.466	22.369	25.494	1.00 27.92	С
	ATOM	978	С	LYS A	294	17.529	23.530	25.786	1.00 27.57	С
	ATOM	979	0	LYS A	294	16.947	24.114	24.866	1.00 27.85	0
	MOTA	980	CB	LYS A		19.618	22.833	24.595	1.00 31.41	С
15	ATOM	981	CG	LYS A		20.500	23.907	25.239	1.00 32.77	С
	ATOM	982	CD	LYS A		21.578	24.416	24.284	1.00 36.06	C
	ATOM	983	CE	LYS A		22.872	23.633	24.419	1.00 37.32	Ċ
	ATOM	984	NZ	LYS A		23.599	23.990	25.673	1.00 37.49	N
	ATOM	985	N	TYR A		17.363	23.852	27.061	1.00 25.07	N
20	ATOM	986	CA	TYR A		16.465	24.928	27.451	1.00 24.97	C
20	ATOM	987	C	TYR A		17.208	26.154	27.938	1.00 25.69	Č
	ATOM	988	Ö	TYR A		18.005	26.074	28.865	1.00 24.37	Ö
	ATOM	989	СВ	TYR A		15.517	24.431	28.543	1.00 24.19	c
	ATOM	990	CG	TYR A		14.927	23.080	28.216	1.00 24.13	C
25	ATOM	991	CD1			15.297	21.943	28.932	1.00 24.03	C
23			CD1			14.023	22.933	27.167	1.00 23.33	C
	MOTA	992	_	TYR A		14.023		28.611	1.00 23.84	C
	MOTA	993			-		20.692		1.00 24.83	C
	ATOM	994	CE2			13.500	21.688	26.836		
20	MOTA	995	CZ	TYR A		13.882	20.573	27.563	1.00 24.74	С
30	MOTA	996	ОН	TYR A		13.369	19.338	27.244	1.00 24.72	0
	MOTA	997	N	ARG A		16.921	27.286	27.302	1.00 27.37	N
	MOTA	998	CA	ARG A		17.532	28.566	27.632	1.00 29.21	С
	MOTA	999	C	ARG A		16.457	29.505	28.177	1.00 28.74	C
0.5	MOTA	1000	0	ARG A		15.269	29.177	28.171	1.00 28.04	0
35	MOTA	1001	СВ	ARG A		18.140	29.201	26.377	1.00 31.67	С
	MOTA	1002	CG	ARG A		19.115	28.332	25.590	1.00 36.71	С
	MOTA	1003	CD	ARG A		19.581	29.091	24.352	1.00 40.42	С
	MOTA	1004	NE	ARG A		20.676	28.444	23.631	1.00 44.23	N
	ATOM	1005	cz	ARG A		20.533	27.442	22.769	1.00 46.02	С
40	ATOM	1006		ARG A		19.329	26.949	22.508	1.00 46.77	N
	ATOM	1007		ARG A		21.597	26.941	22.152	1.00 46.52	N
	ATOM	1008	N		A 297	16.879	30.678	28.634	1.00 29.11	N
	MOTA	1009	CA	VAL A		15.956	31.675	29.167	1.00 30.41	C
	MOTA	1010	С		A 297	14.821	31.972	28.187	1.00 30.24	С
45	MOTA	1011	0		A 297	13.655	32.065	28.582	1.00 29.94	0
	MOTA	1012	CB	VAL A	A 297	16.692	33.005	29.475	1.00 30.71	С
	ATÓM	1013	CG1	. VAL 2	A 297	15.686	34.103	29.797	1.00 33.35	С
	ATOM	1014	CG2	VAL A	A 297	17.646	32.811	30.644	1.00 32.09	С
	MOTA	1015	N	SER A	A 298	15.168	32.115	26.912	1.00 30.44	N
50	MOTA	1016	CA	SER A	A 298	14.185	32.430	25.881	1.00 30.65	С
	MOTA	1017	С		A 298	13.106	31.370	25.714	1.00 30.99	С
	MOTA	1018	0	SER A	A 298	11.986	31.680	25.304	1.00 31.34	0
	ATOM	1019	СВ		A 298	14.884	32.675	24.539	1.00 31.86	C
	ATOM	1020	OG		A 298		31.559	24.143	1.00 33.35	Ö
55	ATOM	1021	N		A 299			26.028	1.00 29.88	N
-	ATOM	1022	CA		A 299			25.912	1.00 29.41	C
	ATOM	1023	C		A 299			27.019	1.00 28.20	č
	ATOM	1023	o		A 299			26.827	1.00 28.75	Ö
	AIUN	1024	•	ADE .		40.200	20.770	20.02/	1.00 20.73	9

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	ATOM	1025		ASP A			13.162	27.679	25.979	1.00 31.09	С
	MOTA	1026		ASP A			14.070	27.435	24.797	1.00 34.22	С
	MOTA	1027	OD1	ASP A	29	9	13.589	27.548	23.651	1.00 34.74	0
_	MOTA	1028	OD2	ASP A	29	9	15.263	27.129	25.013	1.00 36.25	0
5	MOTA	1029	N	VAL A	30	0	11.837	29.631	28.183	1.00 27.61	N
	MOTA	1030	CA	VAL A	30	0	10.923	29.760	29.308	1.00 26.53	C
	MOTA	1031	С	VAL A	30	0	9.948	30.913	29.070	1.00 26.97	С
	MOTA	1032	0	VAL A	A 30	0	8.781	30.835	29.449	1.00 26.32	0
	ATOM	1033	СВ	VAL A	4 30	0	11.703	29.972	30.623	1.00 27.74	С
10	ATOM	1034	CG1	VAL A	3 O	0	10.749	29.958	31.811	1.00 29.57	С
-	ATOM	1035		VAL A			12.757	28.871	30.772	1.00 27.69	C
	ATOM	1036	N	THR A			10.420	31.980	28.432	1.00 26.55	N
	ATOM	1037	CA	THR A			9.539	33.106	28.142	1.00 27.35	C
	ATOM	1038	C	THR A			8.507	32.672	27.100	1.00 27.20	Ċ
15	MOTA	1039	Ö	THR I			7.394	33.188	27.069	1.00 27.20	ŏ
• •	ATOM	1040	СВ	THR A			10.324	34.329	27.617	1.00 27.90	Č
		1041		THR A							o
	MOTA	1041		THR			11.097 11.250	33.956	26.472	1.00 29.74 1.00 29.44	
	MOTA							34.861	28.696		C
20	ATOM	1043	N	LYS .			8.875	31.715	26.250	1.00 26.49	N
20	MOTA	1044	CA	LYS .			7.948	31.225	25.232	1.00 27.28	C
	MOTA	1045	C	LYS .			6.886	30.318	25.847	1.00 27.81	C
	MOTA	1046	0	LYS			5.960	29.874	25.160	1.00 27.95	0
	MOTA	1047	CB	LYS			8.701	30.477	24.130	1.00 28.36	С
0.5	MOTA	1048	CG	LYS			9.496	31.386	23.206	1.00 29.79	С
25	MOTA	1049	CD	LYS			10.203	30.586	22.128	1.00 30.72	C
	MOTA	1050	CE	LYS			11.019	31.482	21.209	1.00 32.93	C
	MOTA	1051	NZ	LYS	A 30	)2	12.121	32.161	21.934	1.00 33.88	N
	MOTA	1052	N	ALA			7.019	30.048	27.143	1.00 26.44	N
	MOTA	1053	CA	ALA	A 30	)3	6.052	29.219	27.847	1.00 27.88	С
30	MOTA	1054	С	ALA	A 30	)3	5.130	30.097	28.692	1.00 28.91	С
	ATOM	1055	0	ALA	A 30	)3	4.310	29.592	29.457	1.00 29.81	0
	ATOM	1056	CB	ALA	A 30	)3	6.771	28.199	28.726	1.00 27.38	С
	MOTA	1057	N	GLY	A 3	04	5.279	31.415	28.564	1.00 29.66	N
	MOTA	1058	CA	GLY	A 30	04	4.423	32.328	29.309	1.00 30.57	С
35	MOTA	1059	С	GLY	A 30	04	4.963	32.961	30.582	1.00 31.32	С
	MOTA	1060	0	GLY			4.257	33.735	31.234	1.00 32.07	0
	ATOM	1061	N	HIS			6.202	32.649	30.948	1.00 31.10	N
	ATOM	1062	CA	HIS	A 3	05	6.797	33.216	32.155	1.00 30.95	C
	ATOM	1063	С	HIS			7.656	34.439	31.853	1.00 31.77	Č
40	ATOM	1064	Ō	HIS			8.138	34.610	30.731	1.00 31.65	0
	ATOM	1065	СВ	HIS			7.628	32.155	32.881	1.00 30.92	Ċ
	ATOM	1066	CG	HIS			6.799	31.128		1.00 30.70	C
	ATOM	1067		HIS			6.017	31.430	34.679	1.00 31.24	N
	ATOM	1068		HIS			6.599	29.812	33.331	1.00 31.47	c
45	MOTA	1069		HIS			5.369	30.346	35.067	1.00 31.89	c
40	MOTA	1070		HIS			5.704	29.351	34.265	1.00 30.48	N
	ATOM	1071	N	SER			7.839	35.290	32.860	1.00 30.40	
		1071	CA	SER			8.624	36.511	32.700	1.00 32.01	N
	MOTA	1072		SER			9.982		33.392		С
FΩ	MOTA		С					36.449		1.00 34.00	С
50	MOTA	1074	0	SER			10.265	35.523	34.154	1.00 33.09	
	MOTA	1075	CB	SER			7.842	37.710	33.235	1.00 34.42	
	MOTA	1076	OG	SER			7.739	37.654	34.645	1.00 37.62	
	MOTA	1077	N	LEU			10.813	37.455	33.125	1.00 34.07	
	MOTA	1078	CA	LEU			12.155	37.537	33.694	1.00 34.93	
55	MOTA	1079	C	LEU			12.172	37.666	35.212	1.00 33.80	
	ATOM	1080	0	LEU			13.180	37.364	35.851	1.00 33.69	
	MOTA	1081	CB	LEU			12.923	38.710	33.068	1.00 36.84	
	MOTA	1082	CG	LEU	A 3	07	13.434	38.527	31.634	1.00 39.29	С

	MOTA	1083	CD1	LEU A	<b>A</b> :	307	12.282	38.235	30.685	1.00	40.58	C
	MOTA	1084	CD2	LEU A	Α :	307	14.168	39.784	31.201	1.00	40.01	С
	MOTA	1085	N	GLU 2	<b>A</b> :	308	11.060	38.110	35.789	1.00	33.38	N
	MOTA	1086	CA	GLU Z	Α :	308	10.963	38.265	37.235	1.00	32.81	C
5	MOTA	1087	С	GLU A	A :	308	11.165	36.913	37.917	1.00	31.88	С
	MOTA	1088	0	GLU 2	A.	308	11.558	36.842	39.078	1.00	30.22	0
	MOTA	1089	CB	GLU 2	A	308	9.603	38.856	37.607	1.00	37.03	С
	MOTA	1090	CG	GLU A	A	308	9.308	40.169	36.888	1.00	42.70	С
	ATOM	1091	CD	GLU .	A	308	7.914	40.707	37.166	1.00	45.49	C
10	MOTA	1092	OE1	GLU .	A	308	7.522	41.696	36.507	1.00	46.94	0
	MOTA	1093	OE2	GLU .	A	308	7.214	40.149	38.040	1.00	47.58	0
	ATOM	1094	N	LEU .	A	309	10.898	35.838	37.182	1.00	29.69	N
	MOTA	1095	CA	LEU	A	309	11.081	34.492	37.714	1.00	29.34	С
	MOTA	1096	C	LEU			12.348	33.872	37.130	1.00	28.31	C
15	ATOM	1097	0	LEU			13.160	33.290	37.848		26.92	0
	ATOM	1098	СВ	LEU			9.882	33.605	37.360		28.48	C
	MOTA	1099	CG	LEU			10.037	32.116	37.700		28.85	C
	MOTA	1100		LEU			10.011	31.931	39.211		29.55	C
	ATOM	1101	CD2				8.919	31.312	37.048		29.07	Ċ
20	ATOM	1102	N	ILE			12.524	34.019	35.822		28.87	N
	ATOM	1103	CA	ILE			13.673	33.428	35.142		30.36	c
	MOTA	1104	C	ILE			15.051	33.907	35.590		30.97	Č
	ATOM	1105	ō	ILE			15.948	33.092	35.808		30.03	ŏ
	ATOM	1106	СВ	ILE			13.552	33.605	33.617		31.31	C
25	ATOM	1107	CG1				12.218	33.023	33.139		32.43	Č
20	ATOM	1108	CG2				14.695	32.884	32.918		32.83	č
	ATOM	1109	CD1				11.920	33.289	31.681		33.95	c
	ATOM	1110	N	GLU			15.240	35.213	35.726		31.83	N
	MOTA	1111	CA	GLU			16.547	35.707	36.151		33.28	C
30	ATOM	1112	C	GLU			16.945	35.175	37.528		31.76	c
30	ATOM	1113	o	GLU			18.067	34.707	37.714		31.24	o
	ATOM	1114	СВ	GLU			16.573	37.237	36.128		35.65	C
	ATOM	1115	CG	GLU			16.550	37.788	34.710		41.13	C
	MOTA	1116	CD	GLU			16.753	39.287	34.649		43.32	c
35	ATOM	1117	OE1				16.858	39.815	33.522		46.68	o
33								39.933	35.718		45.68	
	ATOM	1118	OE2				16.807		38.511		30.94	0
	ATOM	1119 1120	N CA			312 312	16.032	35.232 34.728	39.851		29.89	N C
	ATOM	1121	CA			312	16.358		39.817		28.28	c
40	MOTA						16.570	33.212				
40	MOTA	1122 1123	O CB			312	17.321	32.656	40.619	1.00	28.14	0
	MOTA					312	15.132	35.115	40.675		31.93	
	MOTA	1124	CG			312	14.612		39.962			C
	MOTA	1125	CD			312	14.740	35.943	38.523		31.29 26.85	C
15	MOTA	1126	N			313	15.896	32.550	38.883			N
45	ATOM	1127	CA			313	16.013	31.102	38.739		26.51	C
	ATOM	1128	C			313	17.425	30.764	38.267		25.16	C
	MOTA	1129	0			313	18.063	29.855	38.788		24.33	0
	MOTA	1130	CB			313		30.583	37.715		27.97	C
50	MOTA	1131	CG			313		29.198	37.935		31.36	C
50	ATOM	1132		LEU				28.676	36.600		29.96	С
	MOTA	1133		2 LEU				28.230	38.536		30.03	С
	MOTA	1134	N			314		31.504	37.279		25.12	N
	MOTA	1135	CA			314		31.255	36.763		25.36	C
	MOTA	1136	С			314		31.552	37.839		25.44	
55	ATOM	1137	0			314		30.802	38.008		25.07	
	MOTA	1138	CB			314		32.122	35.517		26.51	
	MOTA	1139		l ILE				31.811	34.400		28.21	
	MOTA	1140	CG:	2 ILE	A	314	20.982	31.844	35.028	1.00	26.75	C

	ATOM	1141		ILE A 31		18.654	30.407	33.858		С
	MOTA	1142		LYS A 31		20.112	32.641	38.574		N
	MOTA	1143		LYS A 31		21.058	32.994	39.626		C
<b>E</b>	MOTA	1144		LYS A 31		21.117	31.869	40.656		С
5	MOTA	1145		LYS A 3		22.193	31.522	41.149		0
	MOTA	1146		LYS A 3		20.651	34.310	40.296		C
	MOTA	1147		LYS A 3		21.759	34.926	41.134		C
	MOTA	1148		LYS A 3		21.562	36.427	41.306		C
40	MOTA	1149	CE	LYS A 3		22.806	37.082	41.891		С
10	ATOM	1150	NZ	LYS A 3		23.154	36.521	43.227	1.00 41.56	N
	ATOM	1151	N	PHE A 3		19.958	31.295	40.967	1.00 23.92	N
	ATOM	1152	CA	PHE A 3		19.874	30.196	41.921	1.00 23.22	C
	MOTA	1153	C	PHE A 3		20.662	28.997	41.400	1.00 22.36	C
15	ATOM	1154	0	PHE A 3		21.422	28.380	42.151	1.00 22.35	0
15	MOTA	1155	CB	PHE A 3		18.410	29.791	42.144	1.00 24.22	C
	MOTA	1156	CG	PHE A 3		18.242	28.546	42.979	1.00 26.30	C
	MOTA	1157		PHE A 3		18.323	28.605	44.370	1.00 27.43	C
	MOTA	1158		PHE A 3		18.037	27.310	42.372	1.00 26.87	C
20	ATOM	1159		PHE A 3		18.204	27.446	45.141	1.00 28.46	C
20	MOTA	1160		PHE A 3		17.918	26.145	43.135	1.00 27.51	C
	MOTA	1161	CZ	PHE A 3		18.002	26.218	44.520	1.00 28.27	C
	MOTA	1162	N	GLN A 3 GLN A 3		20.480	28.665	40.120	1.00 21.28	И
	ATOM	1163	CA	GLN A 3		21.175	27.524 27.681	39.522	1.00 21.35 1.00 21.92	C
25	MOTA	1164	C			22.694 23.410		39.586 39.913		
25	MOTA	1165	O	GLN A 3		20.754	26.735	38.057	1.00 20.68 1.00 21.98	O C
	ATOM	1166 1167	CB CG	GLN A 3		19.296	27.324 26.891	37.855	1.00 21.38	C
	ATOM ATOM	1168	CD	GLN A 3		18.968	25.585	38.563	1.00 22.78	c
	ATOM	1169		GLN A 3		19.792	24.670	38.619	1.00 25.08	0
30	ATOM	1170	NE2			17.756	25.488	39.093	1.00 20.00	N
30	ATOM	1171	NEZ	VAL A 3		23.188	28.870	39.259	1.00 22.14	N
	ATOM	1172	CA	VAL A 3		24.629	29.108	39.301	1.00 22.36	C
	ATOM	1173	C	VAL A 3		25.162	28.983	40.734	1.00 24.71	c
	ATOM	1174	Ö	VAL A 3		26.199	28.349	40.971	1.00 26.38	Ö
35	ATOM	1175	CB	VAL A 3		24.975	30.510	38.727	1.00 24.56	Ċ
00	ATOM	1176		VAL A 3		26.458	30.798	38.897	1.00 26.05	Ċ
	ATOM	1177		VAL A 3		24.608	30.567	37.255	1.00 23.60	Ċ
	ATOM	1178	N	GLY A 3		24.447	29.574	41.687	1.00 25.34	N
	ATOM	1179	CA	GLY A 3		24.868	29.515	43.076	1.00 26.42	C
40	ATOM	1180	C	GLY A 3		24.892	28.099	43.623	1.00 26.70	Č
	ATOM	1181	ō	GLY A 3		25.778	27.738	44.399	1.00 26.15	Ō
	MOTA	1182	N			23.915				N
	ATOM	1183	CA	LEU A 3		23.856	25.910	43.680	1.00 26.49	С
	ATOM	1184	С	LEU A 3		25.001	25.141	43.019	1.00 26.16	С
45	MOTA	1185	0	LEU A 3		25.674	24.342	43.666	1.00 25.62	0
	ATOM	1186	СВ	LEU A		22.499	25.289	43.318	1.00 26.17	С
	MOTA	1187	CG	LEU A		22.202	23.895	43.877	1.00 29.00	С
	MOTA	1188	CD1	LEU A		22.305	23.911	45.394	1.00 28.44	С
	MOTA	1189	CD2	LEU A	320	20.803	23.457	43.439	1.00 27.28	C
50	MOTA	1190	N	LYS A	321	25.231	25.402	41.734	1.00 26.97	N
	MOTA	1191	CA	LYS A	321	26.312	24.743	41.000	1.00 29.33	С
	MOTA	1192	С	LYS A	321	27.664	24.983	41.649	1.00 30.36	С
	ATOM	1193	0	LYS A	321	28.486	24.070	41.746	1.00 30.00	0
	MOTA	1194	СВ	LYS A		26.385	25.252	39.561	1.00 30.33	C
55	ATOM	1195	CG	LYS A	321	25.578	24.465	38.559	1.00 33.36	С
	ATOM	1196	CD	LYS A	321	26.140	23.069	38.341	1.00 33.34	С
	MOTA	1197	CE	LYS A	321	25.279	22.329	37.336	1.00 33.36	С
	ATOM	1198	NZ	LYS A	321	25.668	20.911	37.111	1.00 32.77	N

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	ATOM	1199	N	LYS A	Α :	322	27.894	26.222	42.077	1.00	30.82	N
	MOTA	1200	CA	LYS A	Α :	322	29.155	26.601	42.702	1.00	32.19	С
	MOTA	1201	С	LYS A	A :	322	29.447	25.934	44.037	1.00	32.03	C
	MOTA	1202	0	LYS A	A :	322	30.598	25.896	44.462	1.00	32.87	0
5	ATOM	1203	СВ	LYS A	<b>A</b> :	322	29.234	28.122	42.866	1.00	33.78	C
	ATOM	1204	CG	LYS A	A :	322	29.592	28.853	41.587	1.00	37.24	С
	MOTA	1205	CD	LYS A	A.	322	29.849	30.328	41.856	1.00	39.61	C
	MOTA	1206	CE	LYS 2	Α	322	30.611	30.964	40.712	1.00	41.25	С
	MOTA	1207	NZ	LYS 2	A	322	31.956	30.335	40.544	1.00	43.80	N
10	MOTA	1208	N	LEU 2	Α	323	28.420	25.415	44.703	1.00	30.51	N
	MOTA	1209	CA	LEU 2	A	323	28.627	24.747	45.985	1.00	31.09	С
	MOTA	1210	С	LEU A	Α	323	29.296	23.392	45.774	1.00	31.05	С
	MOTA	1211	0	LEU .	A	323	29.833	22.805	46.715	1.00	31.05	0
	MOTA	1212	СВ	LEU .	Α	323	27.297	24.544	46.719	1.00	30.29	С
15	MOTA	1213	CG	LEU .	Α	323	26.551	25.784	47.220	1.00	31.62	С
	MOTA	1214	CD1	LEU .	Α	323	25.260	25.359	47.904	1.00	30.41	С
	MOTA	1215	CD2	LEU .	Α	323	27.434	26.570	48.180	1.00	31.32	С
	ATOM	1216	N	ASN	Α	324	29.264	22.908	44.535	1.00	30.91	N
	MOTA	1217	CA	ASN	Α	324	29.854	21.619	44.180	1.00	32.42	С
20	MOTA	1218	С	ASN	Α	324	29.466	20.524	45.165	1.00	32.07	С
	MOTA	1219	0	ASN	Α	324	30.323	19.864	45.755	1.00	32.62	0
	MOTA	1220	CB	ASN	Α	324	31.380	21.722	44.110	1.00	36.14	C
	MOTA	1221	CG	ASN	Α	324	31.853	22.576	42.954	1.00	38.53	С
	ATOM	1222	OD1	ASN	Α	324	32.013	23.789	43.087	1.00	43.04	0
25	ATOM	1223	ND2	ASN	Α	324	32.068	21.947	41.805	1.00	40.87	N
	ATOM	1224	N	LEU	Α	325	28.166	20.326	45.333	1.00	29.80	N
	ATOM	1225	CA	LEU	Α	325	27.667	19.320	46.257	1.00	27.98	C
	ATOM	1226	С	LEU	A	325	27.969	17.890	45.836	1.00	27.42	C
	ATOM	1227	0	LEU	A	325	27.984	17.568	44.648	1.00	27.50	0
30	MOTA	1228	CB	LEU	Α	325	26.149	19.454	46.409	1.00	28.15	C
	ATOM	1229	CG	LEU	Α	325	25.592	20.785	46.907	1.00	28.88	C
	MOTA	1230	CD1	LEU	Α	325	24.072	20.701	46.960	1.00	29.23	С
	MOTA	1231	CD2	LEU	Α	325	26.163	21.105	48.276	1.00	28.09	С
	MOTA	1232	N	HIS	Α	326	28.219	17.033	46.821	1.00	26.59	N
35	ATOM	1233	CA	HIS	Α	326	28.430	15.618	46.546	1.00	25.79	C
	ATOM	1234	С	HIS	Α	326	27.003	15.162	46.264	1.00	25.33	C
	MOTA	1235	0	HIS	Α	326	26.052	15.819	46.695	1.00	23.44	0
	ATOM	1236	СВ	HIS	Α	326	28.935	14.882	47.788	1.00	27.17	C
	ATOM	1237	CG	HIS	Α	326	30.303	15.294	48.231	1.00	27.36	С
40	ATOM	1238	ND1	HIS	Α	326	30.942	14.704	49.301	1.00	28.09	N
	ATOM	1239	CD2	HIS	Α	326	31.159	16.222	47.744	1.00	28.85	С
	MOTA	1240	CE1	HIS	Α	326	32.135	15.251	49.453	1.00	28.02	С
	ATOM	1241	NE2	HIS	Α	326	32.292	16.174	48.521	1.00	29.20	N
	MOTA	1242	N	GLU	Α	327	26.839	14.054	45.554	1.00	24.49	N
45	MOTA	1243	CA	GLU	Α	327	25.497	13.569	45.267	1.00	24.94	С
	MOTA	1244	С	GLU	Α	327	24.768	13.297	46.583	1.00	24.29	С
	ATOM	1245	0	GLU	Α	327	23.553	13.498	46.686	1.00	24.42	0
	MOTA	1246	СВ	GLU	Α	327	25.557	12.302	44.409	1.00	27.30	C
	MOTA	1247	CG	GLU	Α	327	24.185	11.755	44.032	1.00	29.69	
50	MOTA	1248	CD	GLU	Α	327	24.247	10.740	42.903	1.00	32.63	
	MOTA	1249		GLU	Α	327		9.771	43.015		31.56	
	MOTA	1250	OE2	GLU	Α	327	23.519	10.915	41.903	1.00	32.79	
	ATOM	1251	N			328		12.858	47.595		22.79	
	ATOM	1252	CA			328		12.576	48.911		23.11	
55	MOTA	1253	С			328		13.822	49.500		23.46	
	ATOM	1254	0			328		13.750	50.086		23.51	
	ATOM	1255	CB			328		12.083	49.877		24.71	
	MOTA	1256	CG			328		10.666	49.607		25.97	

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	MOTA	1257		GLU			27.584	10.591	48.506	1.00		С
	MOTA	1258	OE1	GLU 2	A	328	28.201	9.512	48.356	1.00	29.81	0
	MOTA	1259	OE2	GLU 2	A	328	27.793	11.591	47.789	1.00	27.61	0
	MOTA	1260	N	GLU :	A	329	24.939	14.965	49.349	1.00	22.58	N
5	MOTA	1261	CA	GLU .	A	329	24.406	16.221	49.861	1.00	23.12	С
	MOTA	1262	С	GLU .	Α	329	23.212	16.678	49.026	1.00	22.73	С
	ATOM	1263	0	GLU	A	329	22.236	17.203	49.558	1.00	22.26	0
	ATOM	1264	СВ	GLU .			25.511	17.281	49.856	1.00		С
	ATOM	1265	CG	GLU			26.608	16.943	50.859	1.00		Ċ
10	ATOM	1266	CD	GLU			27.940	17.599	50.554	1.00		C
. •	ATOM	1267	OE1	GLU			28.825	17.532	51.429	1.00		ŏ
	ATOM	1268	OE2	GLU			28.113	18.160	49.454	1.00		ŏ
	MOTA	1269	N	HIS			23.291	16.450	47.721	1.00		Ŋ
	MOTA	1270	CA	HIS			22.225	16.836	46.803	1.00		C
15	ATOM	1271	C	HIS			20.908	16.139	47.150	1.00		C
13							19.863		47.150		22.10	
	ATOM	1272	0	HIS				16.790				0
	MOTA	1273	CB	HIS			22.638	16.494	45.364		24.13	С
	MOTA	1274	CG	HIS			21.648	16.916	44.321		25.22	С
20	MOTA	1275		HIS			21.357	18.237	44.060		25.99	N
20	MOTA	1276		HIS			20.913	16.190	43.444		25.76	C
	MOTA	1277		HIS			20.489	18.307	43.065		26.73	С
	ATOM	1278		HIS			20.203	17.078	42.674		25.08	N
	MOTA	1279	N	VAL			20.955	14.823	47.334		22.22	N
	MOTA	1280	CA	VAL			19.739	14.072	47.642		23.00	С
25	ATOM	1281	С	VAL			19.185	14.382	49.024		22.12	С
	MOTA	1282	0	VAL	A	331	17.968	14.393	49.218	1.00	21.17	0
	MOTA	1283	CB	VAL	A	331	19.952	12.544	47.490	1.00	22.74	С
	MOTA	1284	CG1	VAL	Α	331	20.363	12.233	46.053	1.00	25.60	С
	MOTA	1285	CG2	VAL	A	331	21.008	12.045	48.466	1.00	25.97	C
30	MOTA	1286	N	LEU	A	332	20.067	14.634	49.986	1.00	21.61	N
	MOTA	1287	CA	LEU	Α	332	19.611	14.967	51.327	1.00	21.81	C
	MOTA	1288	С	LEU	Α	332	18.884	16.311	51.301	1.00	21.82	С
	MOTA	1289	0	LEU	Α	332	17.874	16.489	51.976	1.00	22.23	0
	ATOM	1290	CB	LEU	Α	332	20.796	15.020	52.303	1.00	22.40	С
35	MOTA	1291	CG			332	21.262	13.656	52.824		22.71	С
	ATOM	1292		LEU			22.617	13.777	53.516		23.21	C
	MOTA	1293		LEU			20.214	13.112	53.776		23.85	C
	MOTA	1294	N			333	19.389	17.253	50.508		21.45	N
	MOTA	1295	CA			333	18.763	18.569	50.420		22.43	C
40	ATOM	1296	C			333	17.363	18.478	49.808		21.61	Č
	ATOM	1297	ō			333	16.440	19.157	50.259		21.39	ō
	ATOM	1298	СВ			333		19.521	49.599		23.63	C
	ATOM	1299	CG			333	19.221	21.000	49.597	•	26.05	Č
	ATOM	1300		LEU			19.253	21.557	51.014		26.27	Č
45	ATOM	1301		LEU			20.157	21.785	48.703		26.03	C
40		1302	N			334	17.198	17.654	48.776		21.27	
	ATOM	1302	CA			334	15.878	17.513	48.163		20.93	N C
	MOTA		C			334	14.928	16.881	49.171		21.48	C
	MOTA	1304					13.769	17.263				
EΩ	ATOM	1305	O			334		16.648	49.256		21.52	0
50	ATOM	1306	CB			334			46.896		21.53	C
	MOTA	1307	CG			334		17.318	45.719		22.31	C
	MOTA	1308	SD			334		16.343	44.219		24.84	S
	MOTA	1309	CE			334		14.909	44.612		24.19	C
	MOTA	1310	N			335		15.922	49.950		21.64	
55	ATOM	1311	CA			335		15.255	50.949		21.82	
	MOTA	1312	C			335			52.045		22.81	C
	ATOM	1313	0			335		16.248	52.455		22.95	
	MOTA	1314	CB	ALA	. 7	335	15.355	14.070	51.564	1.00	22.44	С

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	ATOM	1315	N	ILE A 336	15.111	17.041	52.517	1.00 21.90	N
	MOTA	1316	CA	ILE A 336	14.827	18.022	53.560	1.00 22.98	С
	MOTA	1317	С	ILE A 336	13.822	19.050	53.038	1.00 23.92	С
_	MOTA	1318	0	ILE A 336	12.949	19.496	53.772	1.00 23.55	0
5	MOTA	1319	CB	ILE A 336	16.129	18.730	54.020	1.00 23.77	С
	MOTA	1320		ILE A 336	17.021	17.724	54.753	1.00 24.24	C
	MOTA	1321	CG2		15.803	19.914	54.936	1.00 25.02	C
	MOTA	1322	CD1		18.445	18.188	54.950	1.00 27.51	С
	MOTA	1323	N	CYS A 337	13.942	19.411	51.765	1.00 22.82	N
10	ATOM	1324	CA	CYS A 337	13.020	20.365	51.166	1.00 23.92	C
	MOTA	1325	C	CYS A 337	11.582	19.846	51.235	1.00 24.00	С
	MOTA	1326	0	CYS A 337	10.665	20.577	51.605	1.00 25.45	0
	MOTA	1327	CB	CYS A 337	13.410	20.622	49.705	1.00 22.95	C
	MOTA	1328	SG	CYS A 337	12.289	21.736	48.817	1.00 25.85	S
15	MOTA	1329	N	ILE A 338	11.393	18.578	50.886	1.00 23.65	N
	MOTA	1330	CA	ILE A 338	10.070	17.957	50.890	1.00 23.56	С
	MOTA	1331	С	ILE A 338	9.457	17.814	52.284	1.00 26.02	С
	MOTA	1332	0	ILE A 338	8.288	18.153	52.501	1.00 25.70	0
	MOTA	1333	CB	ILE A 338	10.126	16.560	50.231	1.00 23.28	C
20	MOTA	1334	CG1	ILE A 338	10.483	16.704	48.746	1.00 22.98	С
	MOTA	1335	CG2	ILE A 338	8.794	15.839	50.396	1.00 24.00	C
	MOTA	1336	CD1	ILE A 338	10.807	15.387	48.057	1.00 22.98	С
	MOTA	1337	N	VAL A 339	10.242	17.305	53.225	1.00 26.29	N
	MOTA	1338	CA	VAL A 339	9.754	17.106	54.584	1.00 29.21	С
25	MOTA	1339	С	VAL A 339	9.971	18.359	55.430	1.00 29.45	С
	MOTA	1340	0	VAL A 339	10.807	18.378	56.333	1.00 30.57	0
	MOTA	1341	CB	VAL A 339	10.461	15.901	55.241	1.00 30.37	C
	MOTA	1342	CG1	VAL A 339	9.751	15.516	56.524	1.00 31.20	С
	MOTA	1343	CG2	VAL A 339	10.479	14.725	54.277	1.00 31.68	С
30	MOTA	1344	N	SER A 340	9.213	19.407	55.122	1.00 30.04	N
	MOTA	1345	CA	SER A 340	9.309	20.676	55.842	1.00 30.94	С
	ATOM	1346	С	SER A 340	8.061	20.868	56.701	1.00 31.69	С
	MOTA	1347	0	SER A 340	6.940	20.841	56.195	1.00 31.64	0
	MOTA	1348	CB	SER A 340	9.438	21.838	54.853	1.00 32.39	С
35	MOTA	1349	OG	SER A 340	10.664	21.773	54.142	1.00 35.78	0
	MOTA	1350	N	PRO A 341	8.243	21.075	58.013	1.00 32.38	N
	MOTA	1351	CA	PRO A 341	7.107	21.263	58.919	1.00 33.82	С
	ATOM	1352	С	PRO A 341	6.344	22.579	58.774	1.00 35.40	С
	MOTA	1353	0	PRO A 341	5.204	22.688	59.232	1.00 36.23	0
40	ATOM	1354	CB	PRO A 341	7.745	21.111	60.298	1.00 33.68	
	MOTA	1355	CG	PRO A 341	9.110	21.675	60.094	1.00 33.78	С
	ATOM	1356	CD	PRO A 341	9.517	21.088	58.754	1.00 32.78	С
	MOTA	1357	N	ASP A 342		23.570	58.131	1.00 36.36	
	MOTA	1358	CA	ASP A 342		24.866	57.981	1.00 37.82	
45	MOTA	1359	С	ASP A 342	5.580	25.094	56.657	1.00 38.30	С
	MOTA	1360	0	ASP A 342	5.655	26.181	56.084	1.00 39.93	0
	MOTA	1361	CB	ASP A 342	7.304	26.001	58.213	1.00 39.62	
	ATOM	1362	CG	ASP A 342		25.987	57.218	1.00 41.16	С
	MOTA	1363	OD1	. ASP A 342	9.185	26.989	57.152	1.00 43.27	0
50	MOTA	1364	OD2	2 ASP A 342	8.597	24.974	56.505	1.00 42.71	0
	MOTA	1365	N	ARG A 343	4.887	24.072	56.170	1.00 37.31	N
	MOTA	1366	CA	ARG A 343	4.123		54.933	1.00 37.10	С
	MOTA	1367	C	ARG A 343	2.683		55.375	1.00 37.98	
	ATOM	1368	0	ARG A 343	2.198		56.273	1.00 38.15	
55	MOTA	1369	СВ	ARG A 343	4.207	22.911	54.103	1.00 35.71	
	MOTA	1370	CG	ARG A 343	5.595	22.543	53.612		
	MOTA	1371	CD	ARG A 343		23.513	52.565		
	ATOM	1372	NE	ARG A 343			51.879		

	MOTA	1373		ARG A 343	8.062	23.612	51.032	1.00 28.95	C
	MOTA	1374		ARG A 343	7.818	24.886	50.751	1.00 27.66	N
	MOTA	1375		ARG A 343	9.097	23.002	50.472	1.00 28.12	N
_	ATOM	1376	N	PRO A 344	1.979	25.367	54.761	1.00 39.05	N
5	MOTA	1377	CA	PRO A 344	0.592	25.579	55.180	1.00 39.38	C
	ATOM	1378	C	PRO A 344	-0.279	24.351	54.924	1.00 38.87	C
	ATOM	1379	0	PRO A 344	-0.205	23.742	53.858	1.00 39.29	0
	MOTA	1380	CB	PRO A 344 PRO A 344	0.167 0.974	26.786	54.348 53.092	1.00 39.90 1.00 40.56	C C
10	ATOM ATOM	1381 1382	CG CD	PRO A 344	2.332	26.624 26.243	53.630	1.00 40.36	C
10	ATOM	1383	N	GLY A 345	-1.085	23.978	55.912	1.00 39.00	N
	MOTA	1384	CA	GLY A 345	-1.965	22.836	55.746	1.00 38.30	C
	ATOM	1385	C	GLY A 345	-1.567	21.544	56.437	1.00 37.94	c
	ATOM	1386	Ö	GLY A 345	-2.386	20.630	56.537	1.00 36.75	ŏ
15	ATOM	1387	N	VAL A 346	-0.328	21.452	56.914	1.00 37.90	N
	ATOM	1388	CA	VAL A 346	0.125	20.234	57.585	1.00 38.39	C
	ATOM	1389	C	VAL A 346	-0.584	20.046	58.922	1.00 39.30	Ċ
	ATOM	1390	0	VAL A 346	-0.832	21.012	59.643	1.00 39.35	0
	ATOM	1391	СВ	VAL A 346	1.654	20.249	57.827	1.00 38.81	C
20	ATOM	1392	CG1	VAL A 346	2.383	20.409	56.503	1.00 37.92	С
	ATOM	1393	CG2	VAL A 346	2.030	21.366	58.784	1.00 38.02	С
	MOTA	1394	N	GLN A 347	-0.905	18.796	59.247	1.00 39.79	N
	ATOM	1395	CA	GLN A 347	-1.597	18.481	60.492	1.00 40.19	C
	MOTA	1396	С	GLN A 347	-0.631	18.135	61.612	1.00 38.86	C
25	MOTA	1397	0	GLN A 347	-0.657	18.758	62.673	1.00 39.58	0
	MOTA	1398	CB	GLN A 347	-2.564	17.314	60.280	1.00 43.00	С
	MOTA	1399	CG	GLN A 347	-3.565	17.531	59.157	1.00 47.50	С
	ATOM	1400	CD	GLN A 347	-4.526	18.678	59.423	1.00 50.27	C
20	MOTA	1401	OE1	GLN A 347	-4.582	19.209	60.535	1.00 52.52	0
30	MOTA	1402	NE2	GLN A 347	-5.283	19.071	58.402	1.00 51.48	N
	ATOM	1403	N	ASP A 348	0.223	17.144	61.380	1.00 36.34	N
	ATOM	1404	CA	ASP A 348	1.181	16.730	62.398	1.00 35.34	C
	MOTA MOTA	1405 1406	C	ASP A 348 ASP A 348	2.568 3.474	17.313 16.622	62.152 61.679	1.00 33.83 1.00 33.85	C
35	MOTA	1407	СВ	ASP A 348	1.257	15.203	62.458	1.00 34.84	c
00	ATOM	1408	CG	ASP A 348	1.947	14.707	63.712	1.00 35.23	C
	ATOM	1409		ASP A 348	1.907	13.488	63.972	1.00 34.74	ŏ
	ATOM	1410		ASP A 348	2.531	15.539	64.437	1.00 34.97	ŏ
	ATOM	1411	N	ALA A 349	2.727	18.587	62.492	1.00 32.41	N
40	ATOM	1412	CA	ALA A 349	3.991	19.286	62.307	1.00 32.51	С
	ATOM	1413	C	ALA A 349	5.122	18.665	63.121	1.00 32.68	
	MOTA	1414		ALA A 349	6.263			1.00 32.47	0
	ATOM	1415	СВ	ALA A 349	3.829	20.753	62.677	1.00 32.86	
	MOTA	1416	N	ALA A 350	4.804	18.206	64.328	1.00 31.95	N
45	MOTA	1417	CA	ALA A 350	5.809	17.602	65.200	1.00 31.15	С
	MOTA	1418	C	ALA A 350	6.458	16.367	64.578	1.00 30.76	С
	MOTA	1419	0	ALA A 350	7.676	16.190	64.655	1.00 30.22	
	MOTA	1420	CB	ALA A 350	5.180	17.240	66.547	1.00 32.37	
	MOTA	1421	N	LEU A 351	5.643		63.972	1.00 30.64	
50	MOTA	1422	CA	LEU A 351	6.150		63.340	1.00 30.92	
	MOTA	1423	С	LEU A 351	7.032		62.156	1.00 30.72	
	MOTA	1424	0	LEU A 351	8.137		61.995	1.00 30.35	
	MOTA	1425	CB	LEU A 351	4.989		62.848	1.00 32.92	
EE	MOTA	1426	CG	LEU A 351	5.214		62.690	1.00 34.73	
55	MOTA	1427		L LEU A 351	4.073				
	ATOM	1428		LEU A 351					
	ATOM	1429 1430		ILE A 352 ILE A 352					
	MOTA	1420	CA	THE W 225	7.282	16.056	60.158	1.00 29.35	C

	MOTA	1431	С	ILE A		8.628	16.646	60.580	1.00 29.73	С
	MOTA	1432	0	ILE A		9.658	16.371	59.959	1.00 30.37	0
	MOTA	1433	CB	ILE A		6.465	17.107	59.362	1.00 29.44	С
_	MOTA	1434		ILE A		5.175	16.463	58.842	1.00 29.58	С
5	MOTA	1435		ILE A		7.290	17.647	58.193	1.00 28.41	С
	MOTA	1436	CD1	ILE A		4.166	17.452	58.284	1.00 29.04	C
	MOTA	1437	N	GLU A	353	8.626	17.445	61.644	1.00 30.02	N
	MOTA	1438	CA	GLU A		9.857	18.058	62.130	1.00 30.56	
	MOTA	1439	С	GLU A		10.845	17.000	62.613	1.00 29.99	
10	MOTA	1440	0	GLU A		12.050	17.147	62.438	1.00 29.97	
	MOTA	1441	CB	GLU A		9.565	19.048	63.266	1.00 32.59	
	MOTA	1442	CG	GLU A		10.755	19.941	63.615	1.00 35.81	
	MOTA	1443	CD	GLU A		10.462	20.922	64.740	1.00 38.69	_
	MOTA	1444	OE1	GLU A		9.381	21.551	64.723	1.00 40.67	
15	MOTA	1445	OE2	GLU A		11.321	21.075	65.637	1.00 40.80	
	MOTA	1446	N		A 354	10.334	15.935	63.223	1.00 29.90	
	MOTA	1447	CA		A 354	11.191	14.861	63.716	1.00 29.96	
	MOTA	1448	С		A 354	11.871	14.191	62.531	1.00 29.90	
	MOTA	1449	0		A 354	13.064	13.904	62.570	1.00 31.40	
20	MOTA	1450	CB		A 354	10.367	13.843	64.491	1.00 30.18	
	MOTA	1451	N		A 355	11.100	13.940	61.478	1.00 29.82	
	MOTA	1452	CA		A 355	11.638	13.314	60.274	1.00 28.57	
	MOTA	1453	С		A 355	12.687	14.220	59.628	1.00 28.03	
	ATOM	1454	0		A 355	13.754	13.756	59.234	1.00 27.72	
25	ATOM	1455	СВ		A 355	10.514	13.022	59.259	1.00 29.38	
	ATOM	1456	CG1		A 355	9.516	12.036	59.872	1.00 30.07	
	MOTA	1457	CG2		A 355	11.101	12.458	57.964	1.00 30.16	
	ATOM	1458	CD1		A 355	8.251	11.849	59.054	1.00 30.83	
	MOTA	1459	N		A 356	12.398	15.515	59.534	1.00 27.45	
30	MOTA	1460	CA		A 356	13.345	16.444	58.925	1.00 28.48	
	MOTA	1461	C		A 356	14.621	16.566	59.754	1.00 29.24	
	ATOM	1462	0		A 356	15.719	16.622	59.202	1.00 27.50	
	MOTA	1463	СВ		A 356	12.718	17.833	58.739	1.00 28.93	
	MOTA	1464	CG		A 356	13.536	18.753	57.823	1.00 29.6	
35	MOTA	1465	CD		A 356	13.064	20.198	57.844	1.00 31.3	
	MOTA	1466	OE1		A 356	12.996	20.823	58.903	1.00 31.5	
	MOTA	1467	NE2		A 356	12.747	20.742	56.667	1.00 30.0	
	MOTA	1468	N		A 357	14.482	16.613	61.078	1.00 29.5	
4.0	ATOM	1469	CA		A 357	15.656	16.724	61.945	1.00 30.4	
40	ATOM	1470	С		A 357	16.610	15.550	61.739	1.00 29.7	
	MOTA	1471	0		A 357	17.827	15.729	61.727	1.00 30.3	
	MOTA	1472	CB		A 357	15.244			1.00 32.8	
	MOTA	1473	CG		A 357	14.665	18.141	63.812	1.00 34.7	
4.5	MOTA	1474			A 357	14.821	19.110	63.040	1.00 36.6	
45	MOTA	1475			A 357	14.065	18.236	64.905	1.00 36.8	
	MOTA	1476	N		A 358	16.059	14.351	61.577	1.00 30.0	
	MOTA	1477	CA		A 358	16.887	13.167	61.368	1.00 30.1	
	MOTA	1478	С		A 358	17.689	13.309	60.073	1.00 29.9	
	MOTA	1479	0		A 358		12.880	59.996	1.00 29.0	
50	MOTA	1480	CB		A 358		11.906	61.323	1.00 31.1	
	MOTA	1481	CG		A 358			61.149	1.00 33.9	
	ATOM	1482	CD		A 358		9.382	61.402	1.00 36.0	
	ATOM	1483	NE		A 358			60.479	1.00 38.0	
EE	MOTA	1484	CZ		A 358		8.358	60.533	1.00 39.3	
55	ATOM	1485			A 358				1.00 39.2	
	MOTA	1486			A 358			59.653	1.00 39.2	
	ATOM	1487	N		A 359			59.061	1.00 28.4	
	MOTA	1488	CA	LEU	A 359	17.735	14.128	57.776	1.00 27.7	6 C

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	MOTA	1489		LEU A			18.757	15.255	57.890		27.89	С
	MOTA	1490		LEU A			19.853	15.171	57.338		27.13	0
	MOTA	1491		LEU A			16.704	14.482	56.697	1.00	27.85	С
_	ATOM	1492		LEU A			15.646	13.421	56.384		27.95	С
5	MOTA	1493		LEU A			14.593	13.994	55.448		28.15	C
	ATOM	1494	CD2	LEU F	<b>A</b> :	359	16.310	12.210	55.758	1.00	28.81	С
	MOTA	1495	N	SER A	<i>Y</i> :	360	18.393	16.312	58.610	1.00	28.62	N
	MOTA	1496	ÇA	SER A	<b>A</b> :	360	19.288	17.448	58.790	1.00	30.04	C
	MOTA	1497	С	SER A	<b>A</b> :	360	20.540	17.046	59.561		30.70	C
10	MOTA	1498	0	SER A	<b>A</b> :	360	21.647	17.454	59.212	1.00	31.03	0
	MOTA	1499	CB	SER A	Α :	360	18.573	18.578	59.534	1.00	32.12	C
	MOTA	1500	OG	SER A	Α :	360	17.496	19.084	58.765	1.00	36.13	0
	ATOM	1501	N	ASN A	Α.	361	20.367	16.251	60.613	1.00	31.34	N
	ATOM	1502	CA	ASN A	Α	361	21.513	15.816	61.405	1.00	31.58	C
15	MOTA	1503	С	ASN A	A	361	22.417	14.921	60.570	1.00	30.77	С
	ATOM	1504	0	ASN A	A	361	23.637	14.935	60.728	1.00	31.06	0
	ATOM	1505	CB	ASN 2	Α	361	21.055	15.083	62.667	1.00	34.28	С
	MOTA	1506	CG	ASN A	A	361	20.328	15.998	63.637	1.00	37.26	С
	MOTA	1507	OD1	ASN .	A	361	20.736	17.139	63.854	1.00	39.61	0
20	MOTA	1508	ND2	ASN .	A	361	19.252	15.497	64.234	1.00	39.64	N
	MOTA	1509	N	THR .	Α	362	21.815	14.146	59.674	1.00	29.26	N
	MOTA	1510	CA	THR .	A	362	22.583	13.270	58.800	1.00	28.14	С
	MOTA	1511	С	THR .	Α	362	23.419	14.135	57.863	1.00	27.56	С
	MOTA	1512	0	THR	Α	362	24.607	13.879	57.654	1.00	27.15	0
25	MOTA	1513	CB	THR	Α	362	21.654	12.371	57.956	1.00	28.47	C
	MOTA	1514	OG1				20.923	11.495	58.823	1.00	28.00	0
	MOTA	1515	CG2	THR	Α	362	22.461	11.548	56.955	1.00	27.60	C
	MOTA	1516	N	LEU	Α	363	22.795	15.167	57.301	1.00	26.97	N
	MOTA	1517	CA	LEU	Α	363	23.493	16.064	56.388		27.40	С
30	MOTA	1518	C	LEU	Α	363	24.623	16.798	57.100		28.23	С
	MOTA	1519	0	LEU	Α	363	25.736	16.884	56.588		27.96	0
	MOTA	1520	CB	LEU			22.519	17.089	55.782		26.59	С
	MOTA	1521	CG	LEU			23.153	18.156	54.882		26.54	С
	MOTA	1522		LEU			23.829	17.495	53.687		26.43	С
35	MOTA	1523	CD2				22.090	19.142	54.417		26.28	С
	MOTA	1524	N	GLN			24.340	17.325	58.286		29.48	N
	MOTA	1525	CA	GLN			25.360	18.054	59.029		31.77	C
	MOTA	1526	С	GLN			26.530	17.140	59.399		30.91	С
4.0	MOTA	1527	0	GLN			27.691	17.539	59.307		30.91	0
40	MOTA	1528	CB	GLN			24.747	18.681	60.283		33.97	С
	MOTA	1529	CG	GLN			25.579	19.812	60.870		39.97	С
	MOTA	1530	CD	GLN			24.749	20.793			41.73	С
	MOTA	1531		GLN				21.785	62.190		45.56	0
4.5	MOTA	1532		GLN				20.523	61.800		43.48	N
45	MOTA	1533	N	THR				15.910	59.799		30.38	N
	MOTA	1534	CA	THR				14.956	60.176		30.54	C
	MOTA	1535	С	THR				14.561	58.965		29.67	C
	MOTA	1536	0			365		14.454			30.84	0
50	MOTA	1537	CB			365		13.687	60.802		30.66	C
50	MOTA	1538		THR				14.045			32.31	0
	ATOM	1539		THR							31.16	C
	MOTA	1540	N			366		14.348			29.11	N
	MOTA	1541	CA			366		13.976			28.71	C
55	MOTA	1542	C			366		15.051			28.75	
55	MOTA	1543	0			366		14.746			29.49	
	MOTA	1544	CB			366		13.708			27.69	
	MOTA	1545				366					27.26	
	MOTA	1546	נטי	LTYR	A	300	28.234	14.421	53.313	1.00	27.40	С

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	ATOM	1547	CD2	TYR A	A 3	366	28.017	12.079	53.759	1.00	27.61	С
	MOTA	1548	CE1	TYR A	A 3	366	28.912	14.144	52.130	1.00	28.23	C
	MOTA	1549		TYR A			28.697	11.790	52.578	1.00	28.00	С
	MOTA	1550	CZ	TYR A	Α 3	366	29.143	12.825	51.770	1.00	28.28	С
5	ATOM	1551	ОН	TYR Z	A 3	366	29.838	12.546	50.615	1.00	28.34	0
	MOTA	1552	N	ILE A	A :	367	28.692	16.310	56.174	1.00	30.76	N
	MOTA	1553	CA	ILE A	A :	367	29.559	17.412	55.762	1.00	32.74	С
	MOTA	1554	C	ILE 2	<b>A</b> :	367	30.823	17.533	56.614	1.00	35.15	С
	MOTA	1555	0	ILE A	A :	367	31.924	17.688	56.086	1.00	35.56	0
10	MOTA	1556	CB	ILE A	A :	367	28.805	18.763	55.807	1.00	32.16	С
	MOTA	1557	CG1	ILE 2	<b>A</b> :	367	27.685	18.764	54.763	1.00	32.04	С
	ATOM	1558	CG2	ILE A	<b>A</b> :	367	29.769	19.915	55.535	1.00	32.16	С
	ATOM	1559	CD1	ILE :	<b>A</b> :	367	26.790	19.977	54.829	1.00	32.66	С
	MOTA	1560	N	ARG .			30.660	17.465	57.930		38.07	N
15	MOTA	1561	CA	ARG .	A :	368	31.794	17.582	58.842		41.56	С
	MOTA	1562	С	ARG .			32.749	16.406	58.711	1.00	43.08	С
	MOTA	1563	0	ARG			33.963	16.558	58.845		43.32	0
	ATOM	1564	СВ	ARG			31.309	17.652	60.289		43.01	C
	ATOM	1565	CG	ARG			30.469	18.868	60.627	1.00		Č
20	ATOM	1566	CD	ARG			30.023	18.814	62.081		50.17	c
	ATOM	1567	NE	ARG			29.222	17.621	62.348		53.52	N
	ATOM	1568	CZ	ARG			28.703	17.315	63.531		54.12	C
	ATOM	1569	NH1	ARG			28.901	18.117	64.570		55.63	N
	MOTA	1570	NH2	ARG			27.983	16.210	63.676		55.17	N
25	ATOM	1571	N	CYS			32.187	15.234	58.440		44.44	N
20	MOTA	1572	CA	CYS			32.964	14.011	58.333		46.85	C
	ATOM	1573	C	CYS			33.501	13.644	56.949		46.74	c
	ATOM	1574	Ö	CYS			34.641	13.198	56.828		46.26	0
	ATOM	1575	СВ	CYS			32.128	12.848	58.881		48.76	c
30		1576	SG	CYS			32.126	11.238	58.816		56.08	S
30	MOTA											
	ATOM	1577	N	ARG			32.700	13.841	55.905		47.07	N
	MOTA	1578	CA	ARG			33.123	13.457	54.558		47.21	C
	MOTA	1579	C	ARG			33.451	14.563	53.559		47.24	C
25	MOTA	1580	0	ARG			34.058	14.292	52.520		46.93	0
35	ATOM	1581	CB	ARG			32.068	12.533	53.940		47.76	C
	MOTA	1582	CG	ARG			31.827	11.248	54.719		49.65	C
	MOTA	1583	CD	ARG			33.034	10.323	54.660		51.30	С
	MOTA	1584	NE	ARG			32.881	9.160	55.532		52.96	N
40	MOTA	1585	CZ	ARG			31.913	8.254	55.420		53.67	С
40	MOTA	1586	NH1				30.999	8.367	54.465		54.23	N
	MOTA	1587		ARG			31.857	7.236	56.268		54.08	N
	MOTA	1588	N	HIS			33.059	15.799	53.845		46.83	N
	MOTA	1589	CA	HIS			33.340	16.879	52.908		47.01	С
	MOTA	1590	С	HIS			34.670	17.554	53.217		47.99	С
45	MOTA	1591	0	HIS			34.809	18.227	54.237		46.99	0
	MOTA	1592	CB	HIS	Α	371	32.225	17.924	52.927	1.00	45.11	С
	MOTA	1593	CG	HIS	Α	371	32.126	18.713	51.659	1.00	44.21	С
	MOTA	1594	ND1	HIS	Α	371	31.086	18.559	50.768	1.00	43.28	N
	ATOM	1595	CD2	HIS	Α	371	32.959	19.629	51.111	1.00	43.81	С
50	ATOM	1596	CE1	HIS	Α	371	31.282	19.346	49.725	1.00	43.79	C
	ATOM	1597	NE2	HIS	A	371	32.412	20.005	49.907	1.00	43.86	N
	ATOM	1598	N	PRO	Α	372	35.665	17.383	52.331	1.00	49.67	N
	ATOM	1599	CA			372		17.972	52.497		51.25	С
	ATOM	1600	С			372		19.480	52.256		52.68	C
55	MOTA	1601	0			372		20.013	51.489		52.39	0
	ATOM	1602	СВ			372		17.208	51.476		51.31	C
	MOTA	1603	CG			372		16.991	50.366		51.05	C
	ATOM	1604	CD			372		16.549	51.115		50.21	c
												_

	MOTA	1605	N	PRO	Α	373	37.961	20.187	52.914	1.00	54.03	N
	MOTA	1606	CA	PRO	Α	373	38.107	21.640	52.777	1.00	55.28	С
•	ATOM	1607	C	PRO	Α	373	38.693	22.028	51.420	1.00	56.14	С
	ATOM	1608	0	PRO	Α	373	39.284	21.197	50.731	1.00	56.44	0
5	ATOM	1609	CB	PRO	A	373	39.036	21.998	53.932	1.00	55.37	С
	ATOM	1610	CG	PRO	A	373	39.925	20.793	54.011	1.00	55.24	С
	MOTA	1611	CD	PRO	Α	373	38.934	19.653	53.885	1.00	54.59	С
	MOTA	1612	N	PRO	Α	374	38.535	23.299	51.017	1.00	56.81	N
	MOTA	1613	CA	PRO	Α	374	37.848	24.368	51.750	1.00	57.49	C
10	ATOM	1614	С	PRO	A	374	36.324	24.301	51.617	1.00	57.83	C
	MOTA	1615	0	PRO	Α	374	35.642	24.299	52.664	1.00	58.58	0
	ATOM	1616	CB	PRO	Α	374	38.431	25.631	51.127	1.00	57.24	С
	MOTA	1617	CG	PRO	Α	374	38.601	25.226	49.698	1.00	57.37	C
	ATOM	1618	CD	PRO	Α	374	39.194	23.834	49.811	1.00	57.05	С
15	MOTA	1619	N	LEU	Α	378	30.279	26.156	57.018	1.00	49.88	N
	MOTA	1620	CA	LEU	Α	378	29.679	27.221	56.220	1.00	45.66	C
	MOTA	1621	С	LEU	Α	378	28.825	26.586	55.127	1.00	41.60	С
	MOTA	1622	0	LEU	Α	378	27.802	27.138	54.723	1.00	38.14	0
	MOTA	1623	CB	LEU	Α	378	30.769	28.092	55.590		53.84	С
20	ATOM	1624	CG	LEU	Α	378	30.382	29.273	54.702	1.00	57.56	С
	ATOM	1625	CD1	LEU	Α	378	29.709	30.349	55.545	1.00	59.64	С
	MOTA	1626	CD2	LEU	A	378	31.634	29.822	54.017		59.59	C
	ATOM	1627	N	LEU	Α	379	29.370	25.200	54.660	1.00	35.28	N
	ATOM	1628	CA	LEU	Α	379	28.529	24.615	53.626		33.21	С
25	ATOM	1629	С	LEU	Α	379	27.095	24.355	54.080		32.24	C
	MOTA	1630	0	LEU			26.157	24.594	53.325		31.09	Ō
	MOTA	1631	СВ	LEU	Α	379	29.151	23.309	53.121		33.21	С
	MOTA	1632	CG	LEU	Α	379	28.379	22.603	52.003		31.83	С
	ATOM	1633	CD1	LEU			28.301	23.508	50.783		33.04	Ċ
30	ATOM	1634		LEU			29.066	21.292	51.651		32.12	C
	ATOM	1635	N			380	26.917	23.869	55.304		31.72	N
	ATOM	1636	CA			380	25.572	23.588	55.792		32.34	C
	ATOM	1637	С			380	24.717	24.852	55.780		32.63	Č
	ATOM	1638	0			380	23.562	24.833	55.339		31.56	ō
35	ATOM	1639	СВ			380	25.611	23.008	57.208		33.03	Ċ
	ATOM	1640	CG			380	24.239	22.659	57.743		34.66	Č
	ATOM	1641	CD1			380	23.486	21.635	57.169		35.98	Č
	MOTA	1642	CD2			380	23.680	23.373	58.800		35.42	C
	ATOM	1643	CE1			380	22.209	21.333	57.636		36.98	Ċ
40	ATOM	1644	CE2			380	22.410	23.080	59.274			Ċ
	ATOM	1645	CZ			380	21.679	22.060	58.688		37.93	Ċ
	MOTA	1646	OH			380	20.420	21.770	59.154		38.77	ō
	ATOM	1647	N			381	25.288	25.950	56.266		32.03	N
	ATOM	1648	CA			381	24.578	27.223	56.304		32.04	C
45	ATOM	1649	C			381	24.190	27.683	54.902		31.59	c
. •	ATOM	1650	ō			381	23.084	28.187	54.693		32.20	ō
	ATOM	1651	СВ			381	25.443	28.287	56.981		32.84	Č
	ATOM	1652	N			382	25.101	27.515	53.948		30.09	N
	ATOM	1653	CA			382	24.849	27.916	52.570		30.96	C
50	ATOM	1654	C			382	23.739	27.083	51.943		30.08	Č
	ATOM	1655	ō			382	22.989	27.575	51.101		30.60	ō
	ATOM	1656	СВ			382	26.121	27.781	51.731		31.98	C
	ATOM	1657	CG			382	27.223	28.757	52.109		34.76	c
	ATOM	1658	CD			382	28.458	28.545	51.254		38.05	
55	ATOM	1659	CE			382	29.559	29.526	51.615		39.31	c
55	MOTA	1660	NZ			382	30.806	29.245	50.845		41.47	N
	ATOM	1661	N			383	23.648	25.819	52.345		29.65	
	MOTA	1662	CA			383	22.621	24.923	51.821		29.41	
	444 OF	1002	Ų.		-		-c. UZI	22.743	JI.021	1.00	٠٧.٩١	C

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	ATOM	1663	С	MET A	A	383	21.253	25.286	52.389	1.00	29.82	C
	MOTA	1664		MET A			20.250	25.271	51.677	1.00	29.12	0
	MOTA	1665	CB	MET A	A	383	22.958	23.468	52.165	1.00	28.17	C
_	MOTA	1666	CG	MET A	A	383	24.130	22.908	51.381	1.00	28.12	C
5	MOTA	1667	SD	MET A	A.	383	24.510	21.186	51.776	1.00		S
	MOTA	1668	CE	MET 2	A	383	23.099	20.338	51.048	1.00	28.89	C
	MOTA	1669	N	ILE A	A	384	21.215	25.612	53.676	1.00	30.76	N
	ATOM	1670	CA	ILE	A	384	19.960	25.983	54.319	1.00	32.84	C
	MOTA	1671	С	ILE .	A	384	19.422	27.271	53.701	1.00	32.96	С
10	MOTA	1672	0	ILE .	Α	384	18.208	27.458	53.594	1.00	32.83	0
	MOTA	1673	CB	ILE .			20.149	26.186	55.842	1.00	34.53	С
	MOTA	1674	CG1	ILE .	Α	384	20.651	24.889	56.482	1.00	36.66	С
	MOTA	1675	CG2	ILE .	A	384	18.834	26.610	56.482	1.00	36.24	С
	MOTA	1676	CD1	ILE .	A	384	19.744	23.691	56.257	1.00	37.66	C
15	MOTA	1677	N	GLN	A	385	20.328	28.153	53.287	1.00	32.82	N
	ATOM	1678	CA	GLN	Α	385	19.931	29.412	52.669	1.00	33.03	C
	MOTA	1679	С	GLN	Α	385	19.288	29.174	51.303	1.00	32.26	С
	MOTA	1680	0	GLN	Α	385	18.382	29.905	50.901	1.00	30.38	0
	ATOM	1681	СВ	GLN	A	385	21.136	30.342	52.515	1.00	35.19	C
20	MOTA	1682	CG	GLN	Α	385	20.839	31.588	51.692	1.00	39.54	C
	MOTA	1683	CD	GLN	Α	385	19.705	32.421	52.270	1.00	41.95	C
	MOTA	1684	OE1	GLN	Α	385	19.024	33.151	51.545	1.00	44.07	0
	ATOM	1685	NE2	GLN	Α	385	19.504	32.324	53.579	1.00	42.90	N
	MOTA	1686	N	LYS	Α	386	19.756	28.152	50.591	1.00	30.96	N
25	MOTA	1687	CA	LYS	Α	386	19.197	27.840	49.282	1.00	30.88	С
	MOTA	1688	С	LYS	Α	386	17.748	27.415	49.447	1.00	29.98	С
	MOTA	1689	0	LYS	Α	386	16.927	27.635	48.558	1.00	29.72	0
	MOTA	1690	CB	LYS	Α	386	19.985	26.719	48.601	1.00	32.29	С
	MOTA	1691	CG	LYS	Α	386	21.430	27.064	48.310	1.00	35.09	C
30	MOTA	1692	CD	LYS	Α	386	21.539	28.305	47.453	1.00	36.92	С
	MOTA	1693	CE	LYS	Α	386	22.997	28.643	47.170	1.00	39.17	С
	ATOM	1694	NZ	LYS	Α	386	23.133	30.008	46.589	1.00	40.63	N
	ATOM	1695	N	LEU	Α	387	17.433	26.804	50.583	1.00	29.25	N
	ATOM	1696	CA	LEU	Α	387	16.064	26.373	50.833	1.00	29.50	С
35	ATOM	1697	С	LEU	Α	387	15.172	27.604	50.982	1.00	29.30	С
	MOTA	1698	0	LEU	Α	387	14.014	27.594	50.572	1.00	27.98	0
	ATOM	1699	CB	LEU	A	387	15.988	25.503	52.091	1.00	30.49	C
	ATOM	1700	CG	LEU	Α	387	16.625	24.111	51.980	1.00	31.01	С
	ATOM	1701	CD1	LEU	Α	387	16.443	23.363	53.289	1.00	33.01	C
40	MOTA	1702	CD2	LEU	A	387	15.985	23.339	50.839	1.00	31.27	С
	MOTA	1703	N	ALA	Α	388	15.714	28.667	51.566	1.00	28.72	N
	MOTA	1704	CA	ALA	Α	388	14.952	29.903	51.735		29.54	С
	MOTA	1705	С	ALA	Α	388	14.757	30.557	50.367		29.67	C
	ATOM	1706	0	ALA	Α	388	13.696	31.121	50.082	1.00	29.63	0
45	ATOM	1707	CB	ALA	Α	388	15.687	30.856	52.679	1.00	30.01	С
	MOTA	1708	N	ASP	Α	389	15.786	30.479	49.524	1.00	29.62	N
	MOTA	1709	CA	ASP	Α	389	15.730	31.044	48.175	1.00	30.08	С
	MOTA	1710	С	ASP	Α	389	14.625	30.360	47.378	1.00	29.78	С
	MOTA	1711	0	ASP	Α	389	13.917	31.000	46.598	1.00	29.18	0
50	MOTA	1712	CB	ASP	Α	389	17.059	30.833	47.445	1.00	31.12	С
	MOTA	1713	CG	ASP	Α	389	18.183	31.682	48.006	1.00	34.91	С
	MOTA	1714	OD1	ASP	Α	389	19.352	31.423	47.643	1.00	36.63	0
	MOTA	1715	OD2	ASP	Α	389	17.901	32.608	48.796		35.52	0
	MOTA	1716	N			390		29.052	47.572		27.64	N
55	ATOM	1717	CA			390		28.262	46.881		28.50	
	MOTA	1718	С	LEU	A	390		28.730	47.223		27.80	
	ATOM	1719	0			390		28.740	46.360		26.98	
	MOTA	1720	CB			390		26.786	47.252		28.87	

	MOTA	1721	CG	LEU A		14.130	25.761	46.221	1.00	31.08	С
	MOTA	1722		LEU A		14.754	26.424	45.017	1.00	31.00	C
	MOTA	1723	CD2	LEU A		15.101	24.810	46.902	1.00	31.75	С
_	MOTA	1724	N	ARG A	391	11.849	29.109	48.481	1.00 2	27.87	N
5	ATOM	1725	CA	ARG A	391	10.535	29.574	48.917	1.00 2	28.52	C
	MOTA	1726	С	ARG A	391	10.132	30.808	48.125	1.00		С
	MOTA	1727	0	ARG A	391	8.968	30.961	47.757	1.00		0
	MOTA	1728	СВ	ARG A		10.536	29.919	50.415	1.00		Ċ
	ATOM	1729	CG	ARG A		10.795	28.744	51.354	1.00		č
10	ATOM	1730	CD	ARG A		9.743	27.658	51.208	1.00		c
. •	ATOM	1731	NE	ARG A		9.952	26.552	52.141	1.00	-	N
	ATOM	1732	CZ	ARG A		9.395	26.460	53.346	1.00		C
	ATOM	1733	NH1	ARG A		8.580	27.411	53.783	1.00		
	ATOM	1734	NH2								N
15						9.646	25.408	54.115	1.00		Ŋ
15	MOTA	1735	N	SER A		11.094	31.690	47.865	1.00		N
	MOTA	1736	CA	SER A		10.811	32.908	47.114	1.00		С
	MOTA	1737	C	SER A		10.483	32.588	45.664	1.00		C
	MOTA	1738	0	SER A		9.577	33.178	45.082	1.00		0
	MOTA	1739	CB	SER A		11.997	33.866	47.185	1.00	-	С
20	ATOM	1740	OG	SER A		12.192	34.305	48.518	1.00	37.19	0
	ATOM	1741	N	LEU A	393	11.219	31.648	45.081	1.00	26.23	N
	MOTA	1742	CA	LEU A	393	10.972	31.253	43.700	1.00	26.10	С
	MOTA	1743	С	LEU A	393	9.614	30.567	43.586	1.00	25.57	С
	MOTA	1744	0	LEU A	393	8.919	30.705	42.576	1.00	26.87	0
25	ATOM	1745	CB	LEU A	393	12.081	30.309	43.216	1.00	26.02	С
	ATOM	1746	CG	LEU A	393	13.450	30.968	43.030		26.66	C
	ATOM	1747	CD1	LEU A		14.536	29.905	42.878	1.00		Ċ
	MOTA	1748		LEU Z		13.400	31.869	41.808		29.45	Č
	ATOM	1749	N		394	9.242	29.825	44.625		24.50	N
30	ATOM	1750	CA		394	7.964	29.122	44.656		26.07	C
•	ATOM	1751	C		394	6.855	30.167	44.570		27.28	C
	ATOM	1752	õ		A 394	5.929	30.055	43.764		26.29	
	ATOM	1753	СВ		394	7.827	28.347	45.967			0
	ATOM	1754	CG		A 394					26.75	C
35						6.646	27.397	45.968		28.26	С
33	ATOM	1755	OD1			5.660	27.604	45.263		28.24	0
	MOTA	1756	ND2			6.736	26.352	46.779		28.79	N
	ATOM	1757	N		A 395	6.966	31.188	45.413		28.62	N
	MOTA	1758	CA		A 395	5.986	32.266	45.464		30.55	С
40	ATOM	1759	С		A 395	5.815	32.976	44.130		29.66	С
40	MOTA	1760	0		A 395	4.691	33.213	43.684		29.50	O
	ATOM	1761	CB		A 395	6.385	33.280	46.536		33.44	С
	ATOM	1762	CG		A 395	6.277	32.744	47.954	1.00	40.01	C
	MOTA	1763	CD	GLU	A 395	4.838	32.481	48.366	1.00	44.38	С
	MOTA	1764	OE1	GLU .	A 395	4.618	32.045	49.518	1.00	46.89	0
45	ATOM	1765	OE2	GLU	A 395	3.924	32.713	47.540	1.00	46.99	0
	ATOM	1766	N	GLU .	A 396	6.929	33.324	43.496	1.00	29.08	N
	MOTA	1767	CA	GLU .	A 396	6.871	34.013	42.217	1.00	28.78	С
	ATOM	1768	С	GLU .	A 396	6.280	33.102	41.148		28.20	C
	MOTA	1769	0	GLU .	A 396	5.486	33.545	40.317		27.96	0
50	MOTA	1770	СВ		A 396	8.265	34.490	41.791		30.45	Č
	ATOM	1771	CG		A 396	8.276	35.254	40.465		30.29	c
	ATOM	1772	CD		A 396	7.502	36.568	40.525		33.32	C
	MOTA	1773		GLU		7.098	37.068	39.452		32.46	
	ATOM	1774		GLU		7.307	37.108				0
55								41.639		32.27	0
55	ATOM	1775	N		A 397	6.651	31.826	41.162		26.94	N
	MOTA	1776	CA		A 397	6.104	30.919	40.162		27.05	C
	MOTA	1777	C		A 397	4.583	30.835	40.295		27.50	C
	MOTA	1778	0	HIS	A 397	3.866	30.834	39.294	1.00	27.05	0

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	ATOM	1779	CB	HIS A	1	397	6.718	29.519	40.282	1.00	26.64	С
	ATOM	1780	CG	HIS A	1	397	6.058	28.507	39.400	1.00	26.04	C
	MOTA	1781	ND1	HIS A	A.	397	4.999	27.731	39.822	1.00	27.22	N
	ATOM	1782	CD2	HIS A	1	397	6.227	28.228	38.086	1.00		C
5	ATOM	1783		HIS A			4.542	27.024	38.805		26.59	Ċ
	ATOM	1784		HIS A			5.268	27.308	37.740		26.30	N
	ATOM	1785	N	SER A			4.094	30.785	41.529		28.04	N
	ATOM	1786	CA	SER A			2.657	30.696	41.775		29.62	C
	ATOM	1787	C	SER A			1.921					
10								31.901	41.195		29.87	C
10	ATOM	1788	0	SER A			0.862	31.761	40.579		28.32	0
	ATOM	1789	СВ	SER A			2.389	30.604	43.279		31.44	С
	MOTA	1790	OG	SER A			1.000	30.483	43.534		38.72	0
	MOTA	1791	N	LYS A			2.485	33.085	41.397	1.00	30.06	N
	MOTA	1792	CA	LYS 2			1.882	34.313	40.885	1.00	31.61	С
15	ATOM	1793	С	LYS A	A	399	1.807	34.283	39.363	1.00	30.91	C
	ATOM	1794	0	LYS A	A	399	0.790	34.651	38.771	1.00	30.21	0
	MOTA	1795	CB	LYS A	A	399	2.698	35.527	41.336	1.00	33.79	С
	ATOM	1796	CG	LYS A	A	399	2.754	35.693	42.842	1.00	38.63	С
	ATOM	1797	CD	LYS A	A	399	3.521	36.946	43.236	1.00	41.48	С
20	ATOM	1798	CE	LYS	A	399	3.571	37.101	44.750		43.57	Č
	ATOM	1799	NZ	LYS		399	4.340	38.313	45.155		44.90	N
	ATOM	1800	N	GLN A			2.886	33.834	38.731		29.20	N
	ATOM	1801	CA	GLN A			2.926	33.770	37.278		28.78	C
	ATOM	1802	C	GLN 3			2.052	32.660	36.702		27.95	c
25	ATOM	1803	0	GLN A			1.524	32.789			27.64	
25				GLN .					35.595			0
	MOTA	1804	CB				4.374	33.637	36.802		28.49	C
	ATOM	1805	CG	GLN .			5.147	34.942	36.964		30.64	С
	ATOM	1806	CD	GLN .			6.483	34.940	36.256		31.34	С
~~	MOTA	1807	OE1				6.673	34.235	35.265		33.90	0
30	ATOM	1808	NE2				7.414	35.751	36.749	1.00	31.02	N
	MOTA	1809	N	TYR	A	401	1.894	31.571	37.446	1.00	26.56	N
	MOTA	1810	CA	TYR	Α	401	1.051	30.481	36.980	1.00	27.31	С
	MOTA	1811	С	TYR	A	401	-0.382	30.998	36.941	1.00	27.98	C
	ATOM	1812	0	TYR	Α	401	-1.147	30.686	36.024	1.00	27.25	0
35	ATOM	1813	CB	TYR	Α	401	1.127	29.285	37.931	1.00	27.75	С
	MOTA	1814	CG	TYR	Α	401	0.229	28.147	37.516	1.00	27.44	С
	ATOM	1815	CD1	TYR	Α	401	0.600	27.281	36.489		28.38	С
	MOTA	1816	CD2				-1.013	27.960	38.119			c
	ATOM	1817	CE1	TYR			-0.242	26.260	36.068	1.00	28.43	Č
40	ATOM	1818	CE2	TYR			-1.868	26.938	37.703		29.62	č
	ATOM	1819	CZ	TYR			-1.475	26.094	36.677	1.00	29.95	c
	ATOM	1820	OH	TYR			-2.319	25.089	36.252	1.00	30.37	0
		1821							37.948			_
	ATOM		N	ARG			-0.742	31.790			29.27	N
15	ATOM	1822	CA	ARG			-2.083	32.360	38.021		32.16	C
45	ATOM	1823	C	ARG			-2.386	33.173	36.769		32.00	C
	MOTA	1824	0	ARG			-3.434	32.998	36.150		31.31	0
	MOTA	1825	СВ	ARG			-2.220	33.251	39.256		36.10	С
	MOTA	1826	CG	ARG			-3.587	33.906	39.391	1.00	41.28	С
	MOTA	1827	CD	ARG			-3.710	34.730	40.669	1.00	45.66	С
50	MOTA	1828	NE	ARG	Α	402	-3.552	33.916	41.873	1.00	49.74	N
	ATOM	1829	CZ	ARG	Α	402	-2.382	33.582	42.410	1.00	51.64	С
	MOTA	1830	NH1	ARG	Α	402	-1.249	33.996	41.856		52.67	N
	ATOM	1831		ARG			-2.343	32.825	43.499		52.69	N
	ATOM	1832	N			403	-1.471	34.066	36.402		32.10	N
55	ATOM	1833	CA	CYS			-1.645	34.895	35.210		33.04	C
-	ATOM	1834	C	CYS			-1.781	34.014	33.976		32.14	C
	ATOM	1835	Õ	CYS			-2.620	34.257	33.106		30.55	o
		1836	СВ			403	-0.450					
	MOTA	1020	CD	CIS	^	403	-0.430	35.838	35.030	1.00	35.60	С

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	MOTA	1837	SG	CYS	A	403	-0.253	36.492	33.340	1.00 4	4.46	S
	MOTA	1838	N	LEU	Α	404	-0.950	32.980	33.911	1.00 3	0.59	N
	MOTA	1839	CA	LEU	A	404	-0.967	32.065	32.784	1.00 3		С
	ATOM	1840	С	LEU	Α	404	-2.327	31.390	32.638	1.00 2		C
5	ATOM	1841	0	LEU	Α	404	-2.840	31.256	31.529	1.00 3		ō
	ATOM	1842	CB	LEU	A	404	0.130	31.008	32.955	1.00 3		Ċ
	ATOM	1843	CG	LEU			0.353	30.078	31.766	1.00 3		C
	ATOM	1844	CD1	LEU			0.840	30.895	30.580		6.26	c
	ATOM	1845		LEU			1.370	29.005	32.127		5.52	C
10	ATOM	1846	N	SER			-2.918	30.987	33.760	1.00 2		N
	ATOM	1847	CA	SER			-4.212	30.309	33.749	1.00 2		C
	MOTA	1848	C	SER			-5.358	31.173	33.218	1.00 2		C
	MOTA	1849	Ö	SER			-6.423	30.651	32.885	1.00 2		
	ATOM	1850	CB	SER			-4.563	29.802	35.153	1.00 2		0
15	ATOM	1851	OG	SER			-4.841	30.873				С
.0	ATOM	1852	N	PHE			-5.147	32.484	36.040	1.00 3		0
	ATOM	1853	CA	PHE					33.145	1.00 2		N
	ATOM	1854	CA	PHE			-6.179	33.396	32.636	1.00 2		С
			0				-6.263	33.340	31.112	1.00 2		С
20	MOTA	1855		PHE			-7.256	33.778	30.518	1.00 2		0
20	ATOM	1856	CB	PHE			-5.868	34.842	33.042	1.00 2		C
	MOTA	1857	CG CD1	PHE			-6.058	35.128	34.503	1.00 2		С
	MOTA	1858		PHE			-5.386	36.196	35.099	1.00 2		С
	MOTA	1859		PHE			-6.920	34.361	35.278	1.00 2		С
25	ATOM	1860		PHE			-5.570	36.494	36.446	1.00 3		С
25	ATOM	1861	CE2	PHE			-7.112	34.651	36.632	1.00 3		С
	MOTA	1862	CZ	PHE			-6.436	35.719	37.214	1.00 3		С
	ATOM	1863	N	GLN			-5.220	32.814	30.478	1.00 2		N
	ATOM	1864	CA	GLN			-5.189	32.748	29.019	1.00 2		С
20	MOTA	1865	C	GLN			-6.155	31.687	28.500	1.00 2		С
30	ATOM	1866	0	GLN			-6.086	30.524	28.903	1.00 2		0
	MOTA	1867	CB	GLN			-3.765	32.448	28.527	1.00 2		C
	ATOM	1868	CG	GLN			-3.571	32.694	27.030		26.23	С
	ATOM	1869	CD	GLN			-3.718	34.165	26.651		26.81	С
25	ATOM	1870	OE1				-4.087	34.494	25.520	1.00 2	28.94	0
35	MOTA	1871	NE2				-3.414	35.052	27.590	1.00 2		N
	ATOM	1872	N			408	-7.083	32.079	27.608	1.00 2	25.83	N
	ATOM	1873	CA			408	-8.052	31.124	27.058	1.00 2		С
	MOTA	1874	С			408	-7.384	29.913	26.398	1.00 2		С
40	MOTA	1875	0			408	-6.389	30.056	25.688	1.00 2	29.12	0
40	ATOM	1876	CB			408	-8.835	31.967	26.054	1.00 2		С
	MOTA	1877	CG			408	-8.824	33.331	26.690	1.00 2		С
	MOTA	1878	CD			408	-7.376			1.00 2	26.30	С
	ATOM	1879	N	GLU	Α	409	-7.941	28.731	26.646	1.00	31.66	N
	MOTA	1880	CA			409	-7.441	27.479	26.078	1.00	34.64	С
45	MOTA	1881	С	GLU	A	409	-6.104	27.014	26.661	1.00	34.56	С
	MOTA	1882	0	GLU	Α	409	-5.480	26.100	26.122	1.00	34.24	0
	MOTA	1883	CB	GLU	Α	409	-7.293	27.606	24.555	1.00	37.88	С
	ATOM	1884	CG	GLU	A	409	-8.511	28.167	23.823	1.00		С
	ATOM	1885	CD	GLU	Α	409	-9.724	27.259	23.887	1.00		С
50	ATOM	1886	OE1	GLU	Α	409	-10.252	27.039	24.998	1.00		0
	ATOM	1887	OE2	GLU	A	409	-10.153	26.766	22.821	1.00		0
	ATOM	1888	N	CYS	Α	410	-5.671	27.628	27.759	1.00		N
	ATOM	1889	CA			410	-4.399	27.267	28.382	1.00		C
	ATOM	1890	C			410	-4.396	25.871	29.002	1.00		Č
55	ATOM	1891	Ö			410	-3.390	25.164	28.943	1.00		Ö
-	ATOM	1892	СВ			410	-4.027	28.299	29.455	1.00		c
	ATOM	1893	SG			410		28.006	30.271	1.00		S
	ATOM	1894	N			411	-5.518	25.472	29.593	1.00		N
					_							7.4

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	ATOM	1895	CA	SER .	A	411	-5.611	24.163	30.235	1.00	35.60	С
	ATOM	1896	С	SER .	A	411	-5.215	23.008	29.319	1.00	35.51	С
	ATOM	1897	0	SER .	Α	411	-4.602	22.040	29.770		35.62	0
	ATOM	1898	CB	SER .	A	411	-7.031	23.928	30.763		36.58	С
5	ATOM	1899	OG	SER	Α	411	-7.959	23.837	29.697	1.00	38.87	0
	ATOM	1900	N	MET	Α	412	-5.561	23.108	28.038		35.70	N
	ATOM	1901	CA	MET	A	412	-5.244	22.053	27.079		36.25	C
	ATOM	1902	С	MET			-3.744	21.912	26.846		34.53	C
	ATOM	1903	0	MET			-3.273	20.867	26.393		34.48	ō
10	ATOM	1904	СВ	MET			-5.936	22.324	25.741		40.58	Ċ
	MOTA	1905	CG	MET			-7.433	22.563	25.850		45.64	Č
	ATOM	1906	SD	MET			-8.214	22.729	24.232		52.62	s
	MOTA	1907	CE	MET			-7.402	24.204	23.610		50.53	č
	MOTA	1908	N	LYS			-2.996	22.965	27.150		31.53	N
15	ATOM	1909	CA	LYS			-1.551	22.944	26.960		30.85	C
	ATOM	1910	C	LYS			-0.831	22.407	28.192		30.52	c
	ATOM	1911	Ō	LYS			0.386	22.236	28.187		30.68	Ö
	ATOM	1912	СВ	LYS			-1.042	24.350	26.632		31.05	C
	MOTA	1913	CG	LYS			-1.557	24.897	25.307		32.36	C
20	ATOM	1914	CD	LYS		-	-1.030	26.296	25.035		32.77	C
	MOTA	1915	CE	LYS			-1.521	26.812	23.689		34.50	C
	ATOM	1916	NZ	LYS			-3.014	26.878	23.622		36.17	N
	ATOM	1917	N	LEU			-1.590	22.142	29.248		30.09	N
	ATOM	1918	CA	LEU			-1.014	21.620	30.484		28.96	C
25	ATOM	1919	C	LEU			-1.393	20.147	30.610		28.33	C
	ATOM	1920	Ö	LEU			-1.654	19.489	29.604		29.10	0
	ATOM	1921	СВ	LEU			-1.544	22.427	31.676		28.64	C
	ATOM	1922	CG	LEU			-1.270	23.934	31.581		30.41	C
	ATOM	1923		LEU			-1.967	24.676	32.711		31.19	
30	ATOM	1924	CD2				0.226	24.179	31.624		30.79	C
00	ATOM	1925	N N	THR			-1.401					C
	MOTA	1926	CA	THR			-1.779	19.624 18.232	31.833		27.70	N
	ATOM	1927	C	THR				18.195	32.071		26.69	C
	ATOM	1928	o	THR			-2.620		33.338	1.00		C
35	ATOM	1929	СВ	THR			-2.548	19.104	34.157		26.39	0
33		1930	OG1				-0.556	17.310	32.307	1.00	26.61	C
	MOTA MOTA	1931	CG2				-0.006	17.570	33.607	1.00		0
		1931				416	0.509	17.546	31.247	1.00	26.48	C
	ATOM ATOM		N				-3.432	17.142	33.516		27.60	N
40		1933	CA			416 416	-4.269	17.037	34.717		27.16	C
40	MOTA	1934	C	_			-3.477	17.169	36.026		27.48	C
	ATOM	1935	0			416	-3.930	17.813	36.975		26.90	0
	ATOM	1936	CB			416			34.564			
	ATOM	1937	CG			416	-5.083	15.555	33.072		28.36	С
15	MOTA	1938	CD			416	-3.752	16.071	32.553		28.22	С
45	ATOM	1939	N			417	-2.294	16.560	36.072		25.90	N
	MOTA	1940	CA			417	-1.460	16.610	37.271		25.39	С
	ATOM	1941	С			417	-0.961	18.031	37.545		24.67	С
	ATOM	1942	0			417	-0.983	18.502	38.685		24.55	0
50	MOTA	1943	CB			417	-0.279	15.643	37.124		25.12	С
50	MOTA	1944	CG			417	0.722	15.507	38.273		25.26	С
	MOTA	1945		. LEU			0.021	15.098	39.564		24.40	С
	MOTA	1946		LEU			1.766	14.470	37.882		25.23	С
	MOTA	1947	N			418	-0.506	18.711	36.500		24.66	N
65	MOTA	1948	CA			418	-0.027	20.080	36.640		25.57	С
55	MOTA	1949	С			418	-1.176	20.971	37.111	1.00	26.33	С
	MOTA	1950	0			418		21.814	37.991	1.00	27.09	0
	MOTA	1951	CB	VAL				20.599	35.297		25.14	C
	ATOM	1952	CG1	. VAL	A	418	0.723	22.112	35.338	1.00	26.89	С

	ATOM	1953		VAL A			1.861	19.912	35.009		25.97	С
	MOTA	1954	N	LEU A	A	419	-2.354	20.769	36.530	1.00	26.33	N
	MOTA	1955	CA	LEU 2	A	419	-3.526	21.556	36.902		27.78	C
_	MOTA	1956	С	LEU 2	A	419	-3.861	21.399	38.382	1.00	29.03	С
5	MOTA	1957	0	LEU 2	A	419	-4.206	22.370	39.052	1.00	30.30	0
	MOTA	1958	CB	LEU !	A	419	-4.733	21.143	36.051	1.00	28.60	С
	MOTA	1959	CG	LEU .			-4.696	21.585	34.586	1.00	30.69	С
	MOTA	1960	CD1	LEU .	Α	419	-5.871	20.975	33.828	1.00	30.94	С
	ATOM	1961	CD2	LEU .	Α	419	-4.743	23.105	34.515	1.00	31.11	С
10	MOTA	1962	N	GLU .	A	420	-3.738	20.184	38.904	1.00	29.76	N
	ATOM	1963	CA	GLU .			-4.056	19.962	40.307	1.00	31.06	С
	ATOM	1964	С	GLU .	Α	420	-3.010	20.514	41.268	1.00	30.59	C
	MOTA	1965	0	GLU	Α	420	-3.344	21.184	42.245	1.00	30.30	0
	MOTA	1966	CB	GLU	Α	420	-4.237	18.478	40.605	1.00	32.62	С
15	MOTA	1967	CG	GLU	Α	420	-4.697	18.251	42.037	1.00	36.69	C
	MOTA	1968	CD	GLU	A	420	-4.267	16.919	42.598	1.00	38.47	С
	ATOM	1969	OE1	GLU	Α	420	-4.631	16.624	43.756	1.00	40.46	0
	MOTA	1970	OE2	GLU	Α	420	-3.561	16.171	41.891	1.00	41.56	0
	MOTA	1971	N	VAL	Α	421	-1.744	20.223	40.992	1.00	31.25	N
20	MOTA	1972	CA	VAL	Α	421	-0.663	20.675	41.855	1.00	32.00	С
	MOTA	1973	С	VAL	Α	421	-0.544	22.191	41.960	1.00	32.63	C
	ATOM	1974	0	VAL	Α	421	-0.355	22.724	43.051	1.00	32.82	0
	MOTA	1975	CB	VAL	A	421	0.694	20.082	41.395	1.00	31.60	C
	MOTA	1976	CG1	VAL	Α	421	1.843	20.676	42.208	1.00	31.31	С
25	ATOM	1977	CG2	VAL	Α	421	0.667	18.567	41.556	1.00	31.20	С
	MOTA	1978	N	PHE	Α	422	-0.670	22.890	40.839	1.00	33.47	N
	MOTA	1979	CA	PHE	A	422	-0.541	24.342	40.857	1.00	34.95	C
	MOTA	1980	С	PHE	A	422	-1.866	25.089	40.872	1.00	35.99	C
	MOTA	1981	0	PHE	Α	422	-1.907	26.284	41.159	1.00	36.37	0
30	MOTA	1982	CB	PHE	Α	422	0.310	24.794	39.670	1.00	34.60	С
	ATOM	1983	CG	PHE	A	422	1.679	24.182	39.656	1.00	34.76	С
	MOTA	1984	CD1	PHE	Α	422	2.093	23.389	38.592	1.00	34.88	С
	MOTA	1985	CD2	PHE	Α	422	2.545	24.369	40.728	1.00	35.16	С
	ATOM	1986	CE1	PHE	A	422	3.348	22.790	38.597	1.00	34.92	С
35	MOTA	1987	CE2	PHE	Α	422	3.801	23.774	40.743	1.00	34.70	С
	MOTA	1988	CZ	PHE	A	422	4.202	22.982	39.674	1.00	34.73	C
	MOTA	1989	N	GLY	A	423	-2.946	24.378	40.570	1.00	37.39	N
	ATOM	1990	CA	GLY	Α	423	-4.261	24.993	40.564	1.00	38.89	С
	MOTA	1991	С	GLY	A	423	-4.914	24.907	41.930	1.00	39.70	С
40	MOTA	1992	0	GLY	Α	423	-5.857	24.099	42.083	1.00	40.52	0
	TER	1993		GLY	A	423						
	HETATM	1994	02	VDX		425	17.029	18.071	34.819	1.00	21.73	0
	HETATM	1995	03	VDX		425	4.489	26.946	35.054	1.00	24.67	0
	HETATM	1996	C1	VDX		425	14.139	17.953	35.755	1.00	20.80	С
45	HETATM	1997	C2	VDX		425	14.879	16.893	34.895	1.00	21.02	С
	HETATM	1998	C3	VDX		425	15.992	17.534	33.962	1.00	21.41	С
	HETATM	1999	C4	VDX		425	15.368	18.672	33.049	1.00	21.29	С
	HETATM	2000	C5	VDX		425	14.622	19.724	33.864	1.00	21.00	С
	HETATM	2001	C6	VDX		425	14.797	21.120	33.792	1.00	20.95	С
50	HETATM	2002	C7	VDX		425	14.174	22.286	34.514	1.00	21.23	
	HETATM	2003	C8	VDX		425	13.966	23.488	34.042	1.00	21.54	
	HETATM	2004	C9	VDX		425	14.354	23.927	32.544		21.77	
	HETATM	2005	C10	VDX		425	13.602	19.075	34.828	1.00	20.74	
	HETATM		C11	XX.		425	13.088	24.490	31.671		21.66	
55	HETATM			VDX		425	12.147	25.443	32.564		22.04	
	HETATM		C13	VDX		425	11.753	24.897	34.070		22.01	
	HETATM			VDX		425		24.538			21.80	
	HETATM	2010	C15	VDX		425	12.661	24.266	36.350		22.22	

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	HETATM 2	011 C1	6 VDX	425	11.429	25.231	36.497	1.00	22.39	С
	HETATM 2	012 C1	.7 VDX	425	11.276	25.934	35.106	1.00		C
	HETATM 2	013 C1	XQV 8.	425	10.769	23.570	33.779	1.00	21.50	С
	HETATM 2	014 C1	.9 VDX	425	12.291	19.455	34.852	1.00	20.77	С
5	HETATM 2	015 C2	XDV 0	425	9.849	26.546	34.726	1.00	22.90	C
	HETATM 2	016 C2	21 VDX	425	9.804	27.956	35.482	1.00	23.65	С
	нетатм 2	017 C2	2 VDX	425	8.575	25.824	35.268	1.00		С
	нетатм 2	018 C2	23 VDX	425	7.331	26.060	34.405	1.00		С
	нетатм 2	019 C2	4 VDX	425	6.152	25.266	34.672	1.00		С
10	нетатм 2	020 C2	25 VDX	425	4.775	25.776	34.336	1.00		С
	нетатм 2	021 C2	26 VDX	425	4.701	26.010	32.842	1.00	25.41	С
	нетатм 2	022 C2	7 VDX	425	3.668	24.730	34.723	1.00		С
	нетатм 2	023 01	L VDX	425	13.119	17.359	36.620	1.00	20.68	0
	нетатм 2	024 0	HOH	500	14.347	10.333	30.796	1.00		0
15	нетатм 2	025 0	нон	501	13.828	12.782	35.922	1.00		0
	нетатм 2	026 0	нон	502	13.846	14.468	42.856	1.00		0
	нетатм 2	027 0	нон	503	19.132	15.890	40.266	1.00		0
	нетатм 2	028 0	HOH	504	15.013	12.029	41.977	1.00		0
	нетатм 2	029 0	HOH	505	13.766	10.118	35.125	1.00		0
20	нетатм 2	030 0	HOH	506	16.290	13.157	34.345	1.00	30.57	0
	нетатм 2	031 0	HOH	507	5.938	22.747	23.179	1.00		0
	нетатм 2	032 0	нон	508	13.771	7.592	35.963	1.00		0
	нетатм 2	033 0	нон	509	12.348	25.386	50.763	1.00		0
	HETATM 2	2034 0	нон	510	28.498	23.703	34.824		37.09	ō
25	нетатм 2	2035 0	нон	511	26.394	10.521	64.086		30.68	ō
	нетатм 2	2036 0	нон	512	20.573	9.150	38.613		30.36	Ō
	нетатм 2	2037 0	нон	513	19.724	30.629	29.203		35.40	ō
	HETATM 2	2038 0	нон	514	4.372	27.504	42.595		31.46	ō
	нетатм 2	2039 0	нон	515	2.808	13.423	33.286		30.93	o
30	нетатм 2	2040 0	нон	516	23.698	20.154	43.135		37.92	Ō
	нетатм 2	2041 0	нон	517	11.325	5.901	37.588		30.12	ō
	нетатм 2	2042 0	нон	518	0.885	13.049	59.537		39.32	O
	HETATM 2	2043 0	нон	519	20.338	11.515	62.065		36.13	O
	HETATM 2	2044 0	нон	520	8.913	6.134	53.451		44.37	ō
35	нетатм 2	2045 0	нон	521	4.924	23.321	44.129		33.51	Ō
	HETATM 2	2046 O	нон	522	16.547	6.409	36.375		32.70	Ō
	HETATM 2	2047 0	нон		8.896	35.918	45.789		45.73	o
	нетатм 2	2048 0	нон	524	26.192	21.542	43.420	1.00	28.56	0
	HETATM 2	2049 O	нон		-5.345	32.214	23.915		35.31	O
40	HETATM 2	2050 O	нон	526	9.488	15.901	22.976		29.33	Ō
	HETATM 2	2051 0	нон	527	5.345	31.465	22.796		31.37	Ō
	HETATM 2	2052 O	нон	528	6.982	20.227	51.589	1.00	32.20	o
	HETATM 2	2053 O	нон	529	4.642	13.886	30.953		31.71	Ō
	HETATM 2	2054 O			-3.764	29.115	25.550		37.63	Ö
45	HETATM 2	2055 O			31.831	9.097	66.550		36.20	ō
	HETATM 2	2056 O			10.178	6.595	32.965		30.94	0
	HETATM 2	2057 0			-1.561	14.197	34.245		33.20	O
	HETATM 2				0.476	12.154	62.160		39.93	ō
	HETATM 2	2059 O			25.970	5.142	53.011		47.31	Ō
50	нетатм 2	2060 O			8.695	5.045	44.801		38.39	0
	HETATM 2				22.396	11.047	39.112		40.45	o
	HETATM 2				13.975	29.983	22.553		36.21	ŏ
	HETATM 2				-6.673	18.195	37.122		36.41	Ö
	HETATM 2				15.926	27.813	55.197		43.43	o
55	HETATM 2				21.922	29.786	26.625		39.42	o
_ •	нетатм 2				29.079	22.924	57.335		43.49	o
	HETATM 2				-8.883	26.986	29.744		47.42	o
	HETATM 2				-2.789	31.232	23.837		38.14	0
							,			_

	нетатм 2	060	0	нон	545	15.578	33.329	45.128	1 00	20 44	_
	HETATM 2		0	нон	546	20.810	2.660	42.920	1.00	59.44 51.44	0
	HETATM 2		0	нон	547	27.448					0
							25.982	58.310		43.04	0
5	HETATM 2		0	НОН	548	21.987	8.152	64.287		43.15	0
3	HETATM 2		0	нон	549	14.435	13.091	64.840		35.87	0
	HETATM 2		0	нон	550	1.276	25.772	21.944		40.66	0
	HETATM 2		0	нон	551	14.102	6.513	31.763		43.70	0
	HETATM 2		0	нон	552	11.990	24.017	53.147		45.62	0
4.0	нетатм 2		0	HOH	553	3.481	24.236	20.666	1.00	35.69	0
10	HETATM 2		0	HOH	554	24.054	13.110	35.770	1.00	37.92	0
	нетатм 2		0	HOH	556	6.857	37.182	44.351	1.00	49.60	0
	нетатм 2	080	0	HOH	557	-8.644	30.901	30.925	1.00	41.21	0
	HETATM 2	081	0	HOH	558	17.767	33.571	43.159	1.00	37.66	0
	нетатм 2	082	0	HOH	559	16.954	26.537	23.238		51.77	0
15	HETATM 2	2083	0	HOH	560	27.386	20.638	40.959		37.25	0
	HETATM 2	2084	0	нон	561	31.418	10.182	50.496		47.27	ŏ
	нетатм 2	2085	0	НОН	562	4.082	21.082	20.610		37.94	ō
	HETATM 2		0	нон	563	14.064	10.706	58.224		42.75	Ö
	нетатм 2		ō	нон	564	23.415	29.835	49.803		45.77	ŏ
20	HETATM 2		ō	НОН	565	14.533	11.393	24.395		36.60	Ö
	HETATM 2		ō	нон	566	-0.868	36.798	40.025		52.17	ŏ
	HETATM 2		ŏ	нон	567	2.865	34.386	33.570		42.56	0
	HETATM 2		ŏ	нон	568	-4.893	19.288	30.751		44.30	
	HETATM 2		0	нон	569	30.643					0
25	HETATM 2		Ö	нон	570	22.702	14.674	61.949		43.28	0
25	HETATM 2		0	нон		13.379	3.372 35.172	47.417		36.93	0
					571			44.109		47.38	0
	HETATM 2		0	нон	572	-1.138	20.698	22.966		53.61	0
	HETATM 2		0	НОН	573	25.589	19.849	33.401		52.13	0
20	HETATM 2		0	НОН	574	23.893	13.360	32.579		45.26	0
30	HETATM 2		0	НОН	575	-7.367	18.485	31.944		48.23	0
	HETATM 2		0	нон	576	2.430	19.200	65.790		45.13	0
	HETATM 2		0	нон	577	20.048	32.028	44.907		46.82	0
	HETATM 2		0	нон	578	20.286	6.713	37.519		43.08	0
0.5	HETATM 2		0	нон	579	25.879	5.448	50.403		48.82	0
35	HETATM 2		0	нон	580	24.905	19.763	39.659	1.00	45.39	0
	HETATM 2		0	HOH	581	2.341	14.233	26.082	1.00	50.76	0
	HETATM 2		0	нон	582	15.248	20.000	60.506	1.00	44.08	0
	HETATM 2		0	HOH	583	22.695	7.038	37.715	1.00	46.55	0
	HETATM 2	2107	0	HOH	584	11.915	16.625	66.479	1.00	52.58	0
40	HETATM 2	2108	0	HOH	585	20.145	35.730	35.936	1.00	46.90	0
	HETATM 2	2109	0	HOH	586	10.735	24.933	16.684	1.00	46.64	0
	HETATM 2	2110	0	HOH	587	1.182	9.495	61.830	1.00	55.88	0
	HETATM 2	2111	0	HOH	588	-3.993	16.527	51.745	1.00	43.33	0
	HETATM 2	2112	0	HOH	589	21.842	29.919	56.624	1.00	42.17	0
45	HETATM 2	2113	0	HOH	590	3.602	25.520	44.494		50.24	0
	HETATM 2	2114	0	нон	591	1.198	23.984	44.777		43.76	0
	HETATM 2	2115	0	нон	592	13.208	27.713	54.123		59.17	0
	HETATM 2		0	нон	593	27.958	7.530	50.434		53.55	0
	HETATM :	2117	0	нон	594	22.594	3.510	64.140		45.66	Ō
50	HETATM :	2118	0	нон	595	30.412	22.979	36.623		71.37	ŏ
	HETATM :		0	нон	596	10.560	15.906	20.574		50.32	ŏ
	HETATM :		ō	нон	597	26.021	3.241	64.667		49.85	0
	HETATM :		Ö	нон	598	19.853	9.062	62.967		56.45	
	HETATM :		Ö	нон	599	12.462	3.992	52.363			0
55	HETATM :		ŏ	нон	600	6.152	35.657			42.46	0
00	HETATM :		0					28.721		46.87	0
	HETATM :			HOH	601	7.626	29.983	53.085		51.73	0
			0	HOH	602	11.547	23.591	57.064		51.07	0
	HETATM :	Z 1 Z Ø	0	нон	603	24.407	19.393	31.035	1.00	53.85	0

	HETATM 212		нон	604	12.538	23.006	18.706	1.00		0
	HETATM 212		HOH	605	1.839	16.469	66.997	1.00	49.40	0
	HETATM 212		HOH	606	1.378	19.964	21.070	1.00		0
_	HETATM 213		HOH	607	5.895	26.935	51.419	1.00		0
5	HETATM 213		HOH	608	13.122	33.698	19.464	1.00	52.90	0
	HETATM 213		HOH	609	27.040	8.636	44.102	1.00	44.22	0
	HETATM 213		HOH	610	18.833	30.775	55.879	1.00		0
	HETATM 213		HOH	611	34.509	17.720	47.771		42.84	0
	HETATM 213		HOH	612	18.356	32.644	25.579	1.00	42.52	0
10	HETATM 213		HOH	613	-2.259	16.235	28.804		56.71	0
	HETATM 213		HOH	614	16.400	38.404	21.700	1.00	46.19	0
	HETATM 213		нон	615	9.340	39.540	19.060		51.44	0
	HETATM 213		HOH	616	20.026	35.074	32.855		47.06	0
	нетатм 214		HOH	617	31.604	8.486	59.428		47.99	0
15	HETATM 214		HOH	618	26.228	8.975	40.708		47.20	0
	HETATM 214		нон	619	0.460	15.378	28.064		50.21	0
	HETATM 214		нон	620	15.771	3.385	48.139		38.09	0
	HETATM 214		HOH	621	25.135	17.914	42.644		60.05	0
	HETATM 214		нон	622	-2.286	29.197	21.618		53.99	0
20	нетатм 214	_	HOH	623	32.865	18.926	45.658		48.11	0
	нетатм 214		нон	624	17.116	13.333	25.240		52.60	0
	нетатм 214		нон	625	-2.809	17.978	56.255		53.36	0
	HETATM 214		нон	626	-3.647	7.885	56.347		63.91	0
or	HETATM 215		HOH	627	17.746	24.596	21.608		59.81	0
25	HETATM 215		нон	628	28.368	5.841	47.861		66.08	0
	HETATM 215		НОН	629	13.641	11.618	66.858		52.02	0
	нетатм 215		нон	630	8.052	20.893	16.742		53.91	0
	HETATM 215		нон	631	8.914	38.015	27.578		56.47	0
00	нетатм 215		нон	632	9.081	13.482	19.627		57.14	0
30	HETATM 215		HOH	633	-4.343	24.969	37.694		51.08	0
	нетатм 215		HOH	634	3.597	28.859	46.576		54.80	0
	HETATM 215		HOH	635	27.905	21.432	28.373		59.49	0
	HETATM 215		нон	636	-4.252	18.337	25.491		47.50	0
0.5	HETATM 216		НОН	637	-2.808	23.046	51.839		49.04	0
35	нетатм 216		нон	638	2.757	25.756	18.437		49.80	0
	HETATM 216		HOH	639	15.470	7.390	63.803		52.42	0
	HETATM 216		нон	640	33.689	11.757	50.784		54.00	0
	HETATM 216		нон	641	6.223	13.352	20.927		49.77	0
. 40	HETATM 216		НОН	642	12.267	32.764	51.605		48.76	0
40	HETATM 216		НОН	644	25.211	3.585	48.391		49.75	0
	HETATM 216		НОН	645	0.619	24.002	51.358		49.46	0
	HETATM 216		НОН	646						
	HETATM 216		НОН	647	0.202	23.805	47.834		52.54	0
15	HETATM 217		нон	648	15.471	8.169	23.816		54.49	0
45	HETATM 217		нон	649	4.098	13.117	28.105		43.97	0
	HETATM 217		нон	650	16.032	4.857	59.064		55.67	0
	HETATM 217		нон	651	~5.591	11.911	55.960		63.35	0
	HETATM 217		нон	652	14.373	4.083	36.218		49.18	0
50	HETATM 217		нон	653	11.138	5.501	59.825		51.19	0
50	HETATM 217		НОН	654	26.262	1.299	50.288		61.20	0
	HETATM 217		HOH	655	4.067	20.751	67.111		51.75	0
	HETATM 217		нон	656	11.291	34.551	23.646		53.35	0
	HETATM 217		НОН	657	2.505	33.743	45.342		58.29	0
55	HETATM 218		нон	658	18.881	-0.886	43.452		60.82	0
55	HETATM 218		HOH	659	-1.930	13.191	62.255		65.05	0
	HETATM 218			660	-3.587	12.153	34.625		51.24	0
	HETATM 218			661	-2.064	26.008	58.110		58.94	0
	HETATM 218	34 O	нон	662	18.842	12.351	64.527	1.00	60.06	0

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HETATM 2	185	0	нон	663	30.991	26.420	51.105	1.00 54.69	0
HETATM 2	186	0	нон	664	16.115	30.354	56.207	1.00 60.96	0
нетатм 2	187	0	HOH	665	36.596	19.242	55.988	1.00 55.83	0

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Table 3

Atomic Structure Coordinate Data of
Polyalanine Model of Conserved VDR LBD

5									<del>_</del>		
3											
	ATOM	1	СВ	PRO	103	-17.052			1.00 78.63	A	С
	ATOM	2	CG	PRO	103	-16.933			1.00 78.57	Α	С
	MOTA	3	С	PRO	103	-15.322			1.00 78.42	A	С
40	MOTA	4	0	PRO	103	-15.845			1.00 78.37	A	0
10	MOTA	5	N	PRO	103	-14.952			1.00 78.63	Α	N
	MOTA	6	CD	PRO	103	-15.422			1.00 78.61	Α	C
	MOTA	7	CA	PRO	103	-15.952			1.00 78.57	Α	С
	MOTA	8	N	VAL	104	-14.202			1.00 78.14	Α	N
	MOTA	9	CA	VAL	104	-13.489			1.00 77.74	Α	С
15	MOTA	10	CB	VAL	104			137.584	1.00 77.77	Α	С
	MOTA	11		VAL	104			138.733	1.00 77.66	Α	С
	MOTA	12		VAL	104	-11.984			1.00 77.68	Α	C
	MOTA	13	С	VAL	104	-14.153	-23.671	136.828	1.00 77.43	Α	C
	MOTA	14	0	VAL	104			136.133	1.00 77.67	Α	0
20	MOTA	15	N	GLN	105			136.636	1.00 76.69	A	N
	MOTA	16	CA	GLN	105	-14.254	-21.567	135.582	1.00 75.70	Α	C
	MOTA	17	CB	GLN	105			135.918	1.00 76.09	Α	С
	MOTA	18	CG	GLN	105			136.067	1.00 76.08	Α	C
	ATOM	19	CD	GLN	105			136.099	1.00 76.03	A	С
25	MOTA	20	OE1	GLN	105			135.107	1.00 75.85	A	0
	ATOM	21	NE2	GLN	105	-11.739	-17.800	137.241	1.00 75.74	Α	N
	ATOM	22	С	GLN	105	-13.637	-21.877	134.223	1.00 74.59	Α	C
	MOTA	23	0	GLN	105	-12.719	-22.691	134.111	1.00 74.90	Α	0
	MOTA	24	N	LEU	106			133.193	1.00 72.98	Α	N
30	MOTA	25	CA	LEU	106	-13.654	-21.381	131.836	1.00 71.07	Α	С
	MOTA	26	СВ	LEU	106	-14.603	-22.279	131.032	1.00 71.27	A	С
	ATOM	27	CG	LEU	106	-14.142	-22.724	129.638	1.00 71.35	Α	С
	ATOM	28		LEU	106	-12.802	-23.437	129.733	1.00 71.22	Α	C
0_	MOTA	29	CD2	LEU	106			129.027	1.00 71.16	Α	C
35	MOTA	30	С	LEU	106	-13.537	-20.002	131.185	1.00 69.48	A	C
	MOTA	31	0	LEU	106			130.693	1.00 69.41	A	0
	MOTA	32	N	SER	107	-12.326	-19.456	131.211	1.00 67.67	Α	N
	ATOM	33	CA	SER	107			130.645	1.00 65.85	Α	C
	MOTA	34	CB	SER	107	-10.516	-18.043	130.383	1.00 65.62	Α	C
40	MOTA	35	OG	SER	107	-10.198	-16.891	129.625	1.00 65.53	Α	0
	MOTA	36	С	SER	107			129.360	1.00 64.86	Α	C
	ATOM	37	0	SER	107			128.573	1.00 64.79	Α	0
	ATOM	38	N	LYS	108			129.154	1.00 63.49	Α	N
	MOTA	39	CA	LYS	108	-13.772	-16.121	127.948	1.00 62.43	Α	C
45	MOTA	40	CB	LYS	108			128.055	1.00 62.56	Α	C
	MOTA	41	CG	LYS	108			128.417	1.00 62.85	Α	C
	MOTA	42	CD	LYS	108			129.776	1.00 63.11	Α	С
	MOTA	43	CE	LYS	108			130.129	1.00 63.62	Α	С
	MOTA	44	NZ	LYS	108			131.449	1.00 63.58	Α	N
50	ATOM	45	C	LYS	108			126.750	1.00 61.42	Α	С
	ATOM	46	0	LYS	108			125.661	1.00 61.44	A	0
	MOTA	47	N	GLU	109			126.959	1.00 60.15	Α	N
	MOTA	48	CA	GLU	109			125.900	1.00 58.91	A	С
	ATOM	49	СВ	GLU	109			126.358	1.00 59.90	A	C
55	MOTA	50	CG	GLU	109	-8.171	-15.661	125.279	1.00 61.96	A	С

	MOTA	51	CD	GLU	109	-6.868	-15.046	125.745	1.00 63.27	A	С
	ATOM	52		GLU	109		-13.866		1.00 64.16	Α	0
	MOTA	53	OE2		109		-15.741		1.00 63.84	Α	0
_	MOTA	54	С	GLU	109	-10.443	-17.682	125.524	1.00 57.30	Α	С
5	MOTA	55	0	GLU	109	-10.154			1.00 56.66	Α	0
	MOTA	56	N	GLN	110	-10.655			1.00 55.60	A	N
	MOTA	57	CA	GLN	110	-10.564			1.00 54.48	Α	С
	MOTA	58	CB	GLN	110	-10.456	-20.723	127.626	1.00 53.38	Α	С
	ATOM	59	CG	GLN	110		-20.512		1.00 52.62	A	С
10	MOTA	60	CD	GLN	110		-21.225		1.00 52.04	Α	С
	MOTA	61	OE1	GLN	110		-21.441		1.00 51.99	Α	0
	MOTA	62	NE2	GLN	110	-10.141	-21.583	130.230	1.00 51.70	Α	N
	MOTA	63	С	GLN	110	-11.754	-20.537	125.503	1.00 54.10	Α	С
	MOTA	64	0	GLN	110	-11.603	-21.426	124.671	1.00 53.77	Α	0
15	ATOM	65	N	GLU	111	-12.938	-20.001	125.772	1.00 53.80	Α	N
	MOTA	66	CA	GLU	111		-20.450		1.00 53.73	A	С
	MOTA	67	CB	GLU	111	-15.389	-19.943	125.774	1.00 54.85	Α	С
	ATOM	68	CG	GLU	111	-15.607	-20.597	127.131	1.00 56.90	A	C
	ATOM	69	CD	GLU	111		-20.172		1.00 58.68	A	C
20	ATOM	70	OE1	GLU	111		-20.349		1.00 60.00	A	o
	MOTA	71	OE2		111		-19.666		1.00 59.57	A	ō
	ATOM	72	С	GLU	111		-20.007		1.00 52.44	Α	č
	ATOM	73	0	GLU	111			122.747	1.00 52.44	A	ŏ
	MOTA	74	N	GLU	112			123.334	1.00 50.93	A	N
25	ATOM	75	CA	GLU	112			121.968	1.00 49.46	A	C
	ATOM	76	СВ	GLU	112			121.956	1.00 50.56	A	Č
	ATOM	77	CG	GLU	112			121.322	1.00 52.74	A	C
	ATOM	78	CD	GLU	112			121.887	1.00 53.94	A	Č
	ATOM	79		GLU	112			123.106	1.00 54.84	A	ō
30	ATOM	80		GLU	112			121.110	1.00 54.87	A	ō
	ATOM	81	С	GLU	112			121.212	1.00 47.53	A	Č
	ATOM	82	O	GLU	112			120.015	1.00 47.52	A	o
	ATOM	83	N	LEU	113			121.922	1.00 45.16	Α	N
	ATOM	84	CA	LEU	113			121.341	1.00 42.62	A	C
35	ATOM	85	CB	LEU	113			122.418	1.00 42.09	A	C
	ATOM	86	CG	LEU	113			122.021	1.00 42.04	A	C
	ATOM	87		LEU	113			123.223	1.00 41.08	A	C
	ATOM	88		LEU	113			120.836	1.00 41.07	A	C
	ATOM	89	C	LEU	113			120.824	1.00 41.07	A	C
40	ATOM	90	Ö	LEU	113			119.670	1.00 39.68	A	0
	ATOM	91	N	ILE	114			121.706	1.00 39.49	A	N
	ATOM	92	CA	ILE	114			121.395	1.00 39.05	A	
	ATOM	93	СВ	ILE	114			122.660	1.00 37.48	A	C
	ATOM	94		ILE	114			122.313	1.00 37.40	A	C
45	ATOM	95		ILE	114			123.709	1.00 37.03	A	C
	ATOM	96		ILE	114			125.034	1.00 35.78	A	C
	ATOM	97	C	ILE	114			120.267	1.00 33.31	A	C
	ATOM	98	Ö	ILE	114			119.440	1.00 39.39		
	ATOM	99	N	ARG	115			120.229	1.00 39.39	A.	0
50	MOTA	100	CA	ARG	115			119.185	1.00 39.79	A	N
00	ATOM	101	CB	ARG	115			119.103	1.00 40.43	A	C
	ATOM	102	CG	ARG	115			119.473	1.00 42.33	A	C
	MOTA	103	CD	ARG	115			118.451		A	C
		103	NE		115				1.00 49.10	A	C
55	MOTA		CZ	ARG				117.470	1.00 51.76	A	N
55	MOTA	105		ARG	115			117.053	1.00 52.78	A	
	MOTA	106		ARG	115			117.207	1.00 53.95	A	
	MOTA	107		ARG	115			116.478	1.00 53.49	A	N
	ATOM	108	С	ARG	115	-14.327	-21.824	117.839	1.00 39.30	Α	С

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	MOTA	109	0	ARG	115	-14.794	-22.357	116.833	1.00 39.26	Α	0
	ATOM	110	N	THR	116	-13.190			1.00 37.95	A	N
	MOTA	111	CA	THR	116	-12.389			1.00 36.97	A	С
_	ATOM	112	CB	THR	116	-11.177			1.00 37.51	Α	С
5	ATOM	113		THR	116	-11.625			1.00 39.12	A	0
	ATOM	114		THR	116	-10.434			1.00 37.41	A	C
	ATOM	115	C	THR	116	-11.887			1.00 35.58	A	С
	ATOM ATOM	116 117	N O	THR LEU	116 117		-22.599 -23.176		1.00 35.61 1.00 33.47	A	0
10	ATOM	118	CA	LEU	117		-24.500		1.00 33.47	A A	С И
10	ATOM	119	CB	LEU	117		-25.143		1.00 31.78	A	C
	MOTA	120	CG	LEU	117			118.426	1.00 30.07	A	C
	ATOM	121	CD1	_	117			119.688	1.00 30.04	A	c
	ATOM	122	CD2		117			117.345	1.00 30.20	A	C
15	ATOM	123	C	LEU	117			116.187	1.00 31.49	A	Ċ
	ATOM	124	Ō	LEU	117			115.195	1.00 31.28	A	ō
	ATOM	125	N	LEU	118			116.876	1.00 31.24	A	N
	ATOM	126	CA	LEU	118			116.487	1.00 30.90	Α	С
	ATOM	127	CB	LEU	118	-15.444	-26.091	117.510	1.00 30.93	Α	С
20	ATOM	128	CG	LEU	118	-15.173	-26.707	118.882	1.00 31.21	A	C
	ATOM	129	CD1	LEU	118	-16.333	-26.391	119.819	1.00 31.43	Α	С
	ATOM	130	CD2	LEU	118			118.737	1.00 30.10	A	С
	MOTA	131	С	LEU	118			115.111	1.00 30.73	A	С
05	ATOM	132	0	LEU	118			114.287	1.00 30.25	Α	0
25	ATOM	133	N	GLY	119			114.872	1.00 30.45	Α	N
	MOTA	134	CA	GLY	119			113.586	1.00 29.84	Α	С
	ATOM	135	C	GLY	119			112.445	1.00 29.41	A	C
	MOTA	136	0	GLY	119			111.411	1.00 29.26	A	0
30	MOTA	137 138	N CA	ALA ALA	120 120			112.634	1.00 28.27	A	N
30	MOTA MOTA	139	CB	ALA	120			111.623 112.006	1.00 27.50 1.00 28.11	A	C
	ATOM	140	C	ALA	120			111.455	1.00 26.73	A A	C
	ATOM	141	Ö	ALA	120			110.336	1.00 26.07	A	0
	MOTA	142	N	HIS	121			112.569	1.00 26.34	A	N
35	MOTA	143	CA	HIS	121			112.542	1.00 25.51	A	C
	ATOM	144	СВ	HIS	121			113.967	1.00 25.42	A	Č
	MOTA	145	CG	HIS	121			114.058	1.00 25.78	A	Č
	ATOM	146	CD2	HIS	121			113.949	1.00 25.53	Α	C
	ATOM	147	ND1	HIS	121			114.240	1.00 26.55	Α	N
40	ATOM	148	CE1	HIS	121	-13.633	-32.398	114.239	1.00 27.01	Α	С
	ATOM	149	NE2	HIS	121			114.064	1.00 27.07	Α	N
	MOTA	150	С	HIS	121			111.857	1.00 25.65	Α	С
	MOTA	151	0	HIS	121			111.000	1.00 23.32	Α	0
45	ATOM	152	N	THR	122			112.233	1.00 26.18	Α	N
45	ATOM	153	CA	THR	122			111.644	1.00 27.73	A	С
	ATOM	154	CB	THR	122			112.310	1.00 27.99	Α	С
	MOTA	155	OG1		122			112.194	1.00 32.40	A	0
	ATOM	156	CG2		122			113.780	1.00 27.74	A	C
50	MOTA	157 158	C O	THR	122 122			110.137 109.368	1.00 27.54	A	C
50	MOTA MOTA	159	Ŋ	THR ARG	123			109.368	1.00 26.85 1.00 28.07	A	0
	ATOM	160	CA	ARG	123			109.713	1.00 28.07	A	N
	MOTA	161	СВ	ARG	123			2 108.141	1.00 29.82	A	C
	ATOM	162	CG	ARG	123			106.141	1.00 29.82	A A	C C
55	ATOM	163	ÇD	ARG	123			106.536	1.00 34.42	A	C
	ATOM	164	NE	ARG	123			107.179	1.00 36.89	A	N
	MOTA	165	CZ	ARG	123			106.714	1.00 36.96	A	
	ATOM	166		ARG	123			3 105.592	1.00 37.38	A	
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	ATOM	167	NH2		123	-17.913			1.00 37.20	Α	N
	MOTA	168	С	ARG	123	-14.628	-28.055	107.415	1.00 28.35	Α	C
	MOTA	169	0	ARG	123	-14.967	-28.431	106.290	1.00 27.61	Α	0
	MOTA	170	N	HIS	124	-13.426	-28.324	107.923	1.00 27.75	A	N
5	ATOM	171	CA	HIS	124	-12.409	-29.016	107.125	1.00 27.66	Α	С
	ATOM	172	СВ	HIS	124	-11.148	-28.147	107.062	1.00 28.26	Α	С
	MOTA	173	CG	HIS	124	-11.395			1.00 29.25	Α	С
	ATOM	174	CD2		124	-11.945			1.00 28.40	A	C
	ATOM	175	ND1		124	-11.081			1.00 29.27	A	N
10	ATOM	176	CE1		124	-11.426			1.00 28.76	Α	C
	ATOM	177	NE2		124	-11.953			1.00 29.33	A	N
	MOTA	178	C	HIS	124			107.478	1.00 26.91	A	C
	ATOM	179	Ö	HIS	124			106.599	1.00 26.66	A	Ö
	ATOM	180	N	MET	125			108.735	1.00 26.00		N
15			CA		125					A	
15	MOTA	181		MET				109.108	1.00 26.25	A	С
	MOTA	182	CB	MET	125			110.025	1.00 26.44	A	С
	MOTA	183	CG	MET	125			109.424	1.00 27.40	A	C
	MOTA	184	SD	MET	125			110.350	1.00 31.87	A	S
00	MOTA	185	CE	MET	125			111.998	1.00 31.61	Α	С
20	MOTA	186	C	MET	125			109.731	1.00 25.47	Α	С
	MOTA	187	0	MET	125			109.446	1.00 25.35	Α	0
	MOTA	188	N	GLY	126			110.581	1.00 24.84	Α	N
	MOTA	189	CA	GLY	126			111.247	1.00 24.24	Α	C
	MOTA	190	C	GLY	126	-15.123	-34.597	110.426	1.00 23.67	Α	С
25	MOTA	191	0	GLY	126	-15.172	-35.739	110.883	1.00 23.66	A	0
	MOTA	192	N	THR	127	-15.581	-34.297	109.215	1.00 22.39	Α	N
	ATOM	193	CA	THR	127	-16.177	-35.339	108.390	1.00 22.17	Α	C
	MOTA	194	CB	THR	127	-17.667	-35.039	108.101	1.00 21.86	Α	C
	ATOM	195	OG1	THR	127	-17.787	-33.751	107.497	1.00 21.82	Α	0
30	ATOM	196	CG2		127			109.387	1.00 22.67	A	С
	ATOM	197	С	THR	127			107.067	1.00 21.87	A	C
	ATOM	198	0	THR	127	-16.065	-36.071	106.118	1.00 21.50	A	0
	ATOM	199	N	MET	128			106.996	1.00 21.50	A	N
	ATOM	200	CA	MET	128			105.746	1.00 21.52	A	C
35	ATOM	201	СВ	MET	128			105.845	1.00 22.11	A	C
••	ATOM	202	CG	MET	128			106.770	1.00 22.14	Α	Ċ
	ATOM	203	SD	MET	128			106.857	1.00 21.82	A	s
	ATOM	204	CE	MET	128			108.065	1.00 22.47	Α	c
	ATOM	205	c	MET	128			105.315	1.00 21.38	A	c
40	ATOM	206	Ö	MET	128			104.131	1.00 21.52	A	Ö
40	ATOM	207	N	PHE	129			106.265	1.00 21.94	A	N
	ATOM	208	CA	PHE	129			105.939	1.00 21.94		
	ATOM	209	СВ	PHE	129			107.219	1.00 21.34	_	
	ATOM	210	CG	PHE	129			107.213	1.00 21.54	A	C
45		211		PHE	129			107.311	1.00 21.07	A	C
40	ATOM	212		PHE	129			107.322	1.00 21.10	A	
	MOTA				129					A	C
	ATOM	213		PHE				107.869	1.00 21.61	A	C
	ATOM	214		PHE	129			109.406	1.00 21.65	A	C
50	ATOM	215	CZ	PHE	129			108.912	1.00 21.57	A	С
50	MOTA	216	С	PHE	129			104.938	1.00 21.50	A	C
	ATOM	217	0	PHE	129			104.197	1.00 20.88	A	0
	MOTA	218	N	GLU	130			104.911	1.00 21.77	A	N
	MOTA	219	CA	GLU	130			103.996	1.00 23.20	Α	C
	ATOM	220		GLU	130			104.298	1.00 23.75	Α	С
55	ATOM	221	CG	GLU	130			105.681	1.00 26.72	Α	C
	MOTA	222	CD	GLU	130			105.968	1.00 27.23	A	С
	MOTA	223		GLU	130			105.154	1.00 27.13	Α	0
	ATOM	224	OE2	GLU	130	-20.321	-37.830	107.014	1.00 28.12	A	0

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	ATOM	225	С	GLU	130	-16.313			1.00 22.81	A	С
	MOTA	226	0	GLU	130	-17.020			1.00 22.98	Α	0
	MOTA	227	N	GLN	131	-15.211	-38.396	102.225	1.00 22.34	Α	N
_	ATOM	228	CA	GLN	131	-14.826			1.00 23.67	Α	C
5	MOTA	229	CB	GLN	131	-14.212	-36.864	100.579	1.00 25.71	Α	С
	MOTA	230	CG	GLN	131	-14.915	-35.665	101.279	1.00 31.91	A	С
	ATOM	231	CD	GLN	131	-16.421	-35.495	100.986	1.00 35.53	Α	С
	MOTA	232	OE1	GLN	131	-17.020			1.00 39.09	Α	0
	ATOM	233	NE2	GLN	131	-17.033			1.00 37.15	A	N
10	ATOM	234	С	GLN	131	-13.871			1.00 22.82	A	C
	MOTA	235	Õ	GLN	131	-13.486		99.186	1.00 22.74	Α	ō
	ATOM	236	N	PHE	132	-13.500			1.00 21.59	A	N
	ATOM	237	CA	PHE	132	-12.585			1.00 20.93	Α	Ĉ
	MOTA	238	CB	PHE	132		-42.289		1.00 19.90	A	c
15	ATOM	239	CG	PHE	132		-41.667		1.00 19.86	A	C
.0	MOTA	240		PHE	132		-40.409		1.00 19.07		c
	MOTA	241		PHE	132		-42.347			A	
	ATOM	242		PHE	132			104.337	1.00 18.20	A	C
									1.00 18.72	A	C
20	MOTA	243		PHE	132			105.371	1.00 18.68	A	C
20	MOTA	244	CZ	PHE	132			105.211	1.00 18.24	A	С
	ATOM	245	C	PHE	132	-13.119		99.658	1.00 20.71	A	С
	ATOM	246	0	PHE	132		-42.675	98.861	1.00 20.20	A	0
	ATOM	247	N	VAL	133		-42.300		1.00 20.69	A	N
	MOTA	248	CA	VAL	133		-43.076		1.00 22.08	Α	С
25	MOTA	249	CB	VAL	133	-16.554	-43.305	98.625	1.00 22.06	A	C
	MOTA	250	CG1	VAL	133	-16.799	-44.205	99.820	1.00 22.30	A	C
	MOTA	251	CG2	VAL	133	-17.281	-41.975	98.794	1.00 20.84	A	C
	MOTA	252	С	VAL	133	-14.825	-42.461	97.056	1.00 23.30	Α	C
	MOTA	253	0	VAL	133	-15.065	-43.110	96.040	1.00 21.98	A	0
30	MOTA	254	N	GLN	134	-14.370	-41.214	97.028	1.00 25.25	Α	N
	MOTA	255	CA	GLN	134	-14.110	-40.505	95.786	1.00 28.42	Α	С
	ATOM	256	СВ	GLN	134		-39.001		1.00 31.35	A	C
	ATOM	257	CG	GLN	134		-38.496		1.00 35.93	A	Ċ
	ATOM	258	CD	GLN	134		-37.018		1.00 38.86	A	Č
35	ATOM	259		GLN	134		-36.224		1.00 40.75	Α	ŏ
	ATOM	260	NE2		134		-36.634		1.00 40.53	A	N
	MOTA	261	C	GLN	134		-40.739		1.00 28.55	A	C
	MOTA	262	Ö	GLN	134		-40.113		1.00 28.59	A	o
	ATOM	263	N	PHE	135		-41.612		1.00 27.76		N
40	MOTA	264	CA	PHE	135		-41.834		1.00 27.76	A	
70	ATOM	265	CB	PHE	135					A	C
		266	CG				-41.361		1.00 27.33	A	C
	ATOM			PHE	135		-39.872		1.00 27.16	A	_
	ATOM	267		PHE	135		-38.978		1.00 27.07	A	C
45	ATOM	268		PHE	135		-39.363		1.00 27.37	A	С
45	ATOM	269		PHE	135		-37.595		1.00 26.82	A	C
	ATOM	270		PHE	135		-37.979		1.00 27.26	Α	С
	MOTA	271	CZ	PHE	135		-37.096		1.00 26.50	Α	С
	MOTA	272	С	PHE	135		-43.256		1.00 26.66	Α	С
	ATOM	273	0	PHE	135		-43.843		1.00 26.40	Α	0
50	MOTA	274	N	ARG	136		-43.784		1.00 26.15	Α	N
	MOTA	275	CA	ARG	136		-45.129		1.00 25.96	Α	С
	ATOM	276	CB	ARG	136	-9.900	-45.115	92.452	1.00 26.56	Α	С
	MOTA	277	CG	ARG	136		-44.063		1.00 29.01	Α	C
	ATOM	278	CD	ARG	136		-43.086		1.00 31.35	A	C
55	MOTA	279	NE	ARG	136		-41.743		1.00 34.32	A	N
	ATOM	280	CZ	ARG	136	-8.939	-40.621		1.00 35.38	A	c
	MOTA	281		ARG	136		-40.660		1.00 34.41	A	N
	ATOM	282		ARG	136		-39.452		1.00 37.45	A	N
				_							7.4

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	MOTA	283	С	ARG	136	-10.530	-46.179	94.623	1.00 25.24	A	С
	MOTA	284	0	ARG	136		-46.819		1.00 24.38	A	0
	MOTA	285	N	PRO	137		-46.371		1.00 24.16		
	MOTA	286	CD	PRO	137		-45.707		1.00 23.61	A	N
5	MOTA	287	CA	PRO	137		-47.366			A	C
	MOTA	288	СВ	PRO	137		-47.081		1.00 23.41	A	C
	ATOM	289	CG	PRO	137		-46.654		1.00 23.30	A	C
	ATOM	290	c	PRO	137		-48.776		1.00 23.80	Α	С
	ATOM	291	ŏ	PRO	137		-49.115		1.00 23.31	Α	С
10	ATOM	292	N	PRO	138		-49.620		1.00 22.63	Α	0
	ATOM	293	CD	PRO	138				1.00 23.12	Α	N
	ATOM	294	CA	PRO	138		-49.369		1.00 23.19	Α	С
	ATOM	295	CB	PRO	138		-50.993		1.00 23.26	Α	C
	ATOM	296	CG				-51.679		1.00 23.98	A	С
15	ATOM	297	C	PRO	138		-50.578	_	1.00 24.11	Α	С
10	ATOM			PRO	138		-51.547		1.00 22.75	Α	С
		298	0	PRO	138		-51.142		1.00 22.46	Α	0
	ATOM	299	N	ALA	139		-52.468	_	1.00 21.31	Α	N
	ATOM	300	CA	ALA	139		-53.061		1.00 21.53	Α	С
20	ATOM	301	СВ	ALA	139		-54.024		1.00 21.98	A	Ċ
20	ATOM	302	С	ALA	139		-53.774		1.00 21.59	A	Ċ
	ATOM	303	0	ALA	139		-53.750	98.282	1.00 20.89	A	ō
	MOTA	304	N	HIS	140		-54.405		1.00 21.14	A	Ŋ
	ATOM	305	CA	HIS	140	-12.253	-55.107	99.199	1.00 21.89	Α	C
0.5	MOTA	306	CB	HIS	140	-10.941	-55.903	99.342	1.00 22.63	A	C
25	ATOM	307	CG	HIS	140	-9.759	-55.062	99.725	1.00 21.19	A	C
	ATOM	308	CD2	HIS	140			100.936	1.00 21.22	A	C
	ATOM	309	ND1	HIS	140	-9.024	-54.347	98.804	1.00 20.38	A	N
	ATOM	310	CE1	HIS	140		-53.650		1.00 21.24	A	
	ATOM	311	NE2	HIS	140			100.726	1.00 22.01		C
30	MOTA	312	С	HIS	140	-12.388	-54.153	100.392	1.00 22.01	A	N
	ATOM	313	0	HIS	140	-12.605	-54.586	101.518	1.00 22.47	A	C
	ATOM	314	N	LEU	141	-12.251	-52 857	100.134		A	0
	MOTA	315	CA	LEU	141	~12.364	-51 827	101.166	1.00 23.58	A	N
	MOTA	316	СВ	LEU	141	-11 777	-50 520	100.634	1.00 23.85	A	С
35	ATOM	317	CG	LEU	141	~10 527	_49 927	100.634	1.00 23.18	A	C
	ATOM	318		LEU	141	-9 667	-51 037	101.294	1.00 24.27	Α	C
	ATOM	319		LEU	141	-9 766	-49 121	101.903	1.00 22.26	Α	С
	ATOM	320	C	LEU	141	_13 912	-49.121	100.262	1.00 21.25	A	С
	ATOM	321	ō	LEU	141	-14 066	51.399	101.603	1.00 24.27	Α	C
40	ATOM	322	N	PHE	142	_14.000	-51.148	102.718	1.00 23.20	Α	0
	ATOM	323	CA	PHE	142	-16 172	-51.902	100.719	1.00 24.16	A	N
	ATOM	324	CB	PHE	142	~17 017	-31.717	101.032	1.00 25.36	A	С
	ATOM	325	CG	PHE	142	-16.898	-51.7/3	99.752	1.00 23.10	Α	С
	ATOM	326		PHE	142			98.901	1.00 22.13	Α	С
45	ATOM	327	CD2		142	-17.570		99.240	1.00 22.32	Α	С
. •	ATOM	328	CE1			-16.087		97.780	1.00 22.24	Α	C
	ATOM	329	CE2		142	-17.432		98.467	1.00 22.25	A	С
	MOTA				142	-15.944		97.006	1.00 22.18	A	С
	ATOM	330 331	CZ	PHE	142	-16.615	-48.242	97.349	1.00 20.96	Α	С
50			C	PHE	142	-16.666	-52.771	102.005	1.00 26.65	Α	C
50	MOTA	332	0	PHE	142	-16.213	-53.914	101.976	1.00 26.20	Α	0
	ATOM	333	N	ILE	143	-17.594	-52.380	102.873	1.00 28.79	Α	N
	ATOM	334	CA	ILE	143	-18.165	~53.310	103.834	1.00 31.69	A	C
	ATOM	335	CB	ILE	143	-19.247	-52.630	104.743	1.00 32.78	A	Ċ
65	ATOM	336	CG2		143	-18.682	-51.372	105.382	1.00 33.71	A	C
55	ATOM	337	CG1		143	-20.516	-52.300	103.943	1.00 33.80	A	Ċ
	ATOM	338	CD1		143	-20.373	-51.225	102.876	1.00 35.73	A	Č
	ATOM	339	C	ILE	143	-18.814	-54.449	103.039	1.00 32.40	A	C
	MOTA	340	0	ILE	143	-19.161	-54.277	101.870	1.00 31.97	A	ō
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		241				40.055		100 500			
	MOTA	341	N	HIS	144	-18.967			1.00 33.54	Α	N
	ATOM	342	CA	HIS	144	-19.568			1.00 35.74	A	С
	MOTA	343	CB	HIS	144	-20.924			1.00 36.54	Α	С
_	MOTA	344	CG	HIS	144	-21.853			1.00 37.08	A	С
5	ATOM	345	CD2		144	-22.508			1.00 36.90	Α	С
	MOTA	346	ND1		144	-22.207	-56.189	104.563	1.00 37.27	Α	N
	MOTA	347	CE1	HIS	144	-23.037	-55.349	105.156	1.00 37.23	Α	С
	ATOM	348	NE2	HIS	144	-23.235	-54.321	104.350	1.00 37.35	Α	N
	ATOM	349	С	HIS	144	-18.648	-57.317	101.932	1.00 36.21	A	С
10	ATOM	350	0	HIS	144		-57.751		1.00 36.04	A	0
	MOTA	351	N	HIS	145		-57.289		1.00 37.47	A	N
	ATOM	352	CA	HIS	145		-57.778		1.00 38.79	A	c
	ATOM	353	СВ	HIS	145		-56.611		1.00 38.32	A	Ċ
	ATOM	354	CG	HIS	145	-16.612		99.379	1.00 38.92	A	C
15	ATOM	355		HIS	145	-17.687		99.416	1.00 38.32	A	c
	ATOM	356		HIS	145	-16.436		98.056	1.00 38.33		
	ATOM	357		HIS	145	-17.365				A	N
								97.325	1.00 38.59	A	С
	ATOM	358		HIS	145	-18.138		98.125	1.00 39.24	A	N
20	MOTA	359	C	HIS	145		-58.564		1.00 39.68	Α	С
20	ATOM	360	0	HIS	145		-58.406		1.00 40.45	A	0
	MOTA	361	N	GLN	146		-59.417		1.00 40.62	Α	N
	ATOM	362	CA	GLN	146		-60.232		1.00 40.83	Α	С
	ATOM	363	CB	GLN	146		-61.585	100.899	1.00 42.96	Α	С
	MOTA	364	CG	GLN	146	-13.528	-61.487	99.376	1.00 46.15	Α	С
25	MOTA	365	CD	GLN	146	-13.498	-62.850	98.685	1.00 48.68	Α	С
	MOTA	366	OE1	GLN	146	-14.422	-63.659	98.824	1.00 49.88	Α	0
	ATOM	367	NE2	GLN	146	-12.430	-63.105	97.934	1.00 49.37	Α	N
	ATOM	368	С	GLN	146	-12.193	-59.464	101.412	1.00 39.30	Α	С
	ATOM	369	0	GLN	146			100.467	1.00 39.67	A	ō
30	MOTA	370	N	PRO	147			102.292	1.00 37.59	A	N
	ATOM	371	CD	PRO	147			103.411	1.00 37.45	A	C
	ATOM	372	CA	PRO	147			102.165	1.00 37.43	A	Ċ
	ATOM	373	СВ	PRO	147			103.367	1.00 36.50	A	C
	ATOM	374	CG	PRO	147			103.580	1.00 30.30		C
35	ATOM	375	C	PRO	147			100.835		A	
55		376		PRO				100.835	1.00 33.38	A	C
	MOTA		0		147				1.00 33.33	A	0
	ATOM	377	N	LEU	148			100.535	1.00 30.98	A	N
	ATOM	378	CA	LEU	148		-58.458	99.305	1.00 29.30	A	C
40	ATOM	379	CB	LEU	148		-57.465		1.00 29.16	Α	С
40	MOTA	380	CG	LEU	148		-56.372	98.263	1.00 29.82	Α	С
	ATOM	381		LEU	148		-55.910	98.289	1.00 29.71	Α	С
	MOTA	382		LEU	148		-56.870	96.883	1.00 29.27	Α	С
	ATOM	383	С	LEU	148		-59.865	99.153	1.00 27.66	Α	С
	ATOM	384	0	LEU	148		-60.406	100.085	1.00 26.44	Α	0
45	ATOM	385	N	PRO	149	-7.057	-60.481	97.977	1.00 26.63	Α	N
	ATOM	386	CD	PRO	149	-7.955	-60.093	96.877	1.00 26.30	Α	С
	ATOM	387	CA	PRO	149	-6.523	-61.827	97.767	1.00 26.49	Α	С
	ATOM	388	CB	PRO	149	-7.042	-62.192	96.376	1.00 26.48	Α	С
	ATOM	389	CG	PRO	149		-61.433	96.288	1.00 26.76	Α	C
50	ATOM	390	С	PRO	149	-4.993	-61.798		1.00 26.14	A	Č
-	ATOM	391	0	PRO	149		-60.749		1.00 26.17	A	ō
	MOTA	392	N	THR	150		-62.961		1.00 26.06	A	N
	ATOM	393	CA	THR	150		-63.130		1.00 25.41		
	ATOM	394	CB	THR	150		-64.632		1.00 25.41	A	C
<b>55</b>		395	OG1		150					A	C
55	ATOM	396					-65.037		1.00 23.28	A	0
	ATOM		CG2		150		-64.888		1.00 25.10	A	С
	MOTA	397	C	THR	150		-62.610		1.00 26.17	A	С
	MOTA	398	0	THR	150	-1.155	-61.933	97.123	1.00 25.67	A	0

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	MOTA	399	N	LEU	151	-2.652	-62.924	95.766	1.00 26.09	A	N
	ATOM	400	CA	LEU	151	-1.973	-62.520	94.543	1.00 27.02	Α	С
	MOTA	401	CB	LEU	151	-1.976		93.548	1.00 28.10	Α	С
_	MOTA	402	CG	LEU	151	-0.752		93.469	1.00 28.68	Α	С
5	ATOM	403	CD1		151		-64.874	94.830	1.00 28.89	Α	С
	MOTA	404	CD2		151		-65.927	92.809	1.00 28.50	Α	C
	MOTA	405	С	LEU	151	-2.516		93.865	1.00 26.77	Α	С
	MOTA	406	0	LEU	151		-60.827	92.849	1.00 26.56	Α	0
4.0	MOTA	407	N	ALA	152		-60.659	94.422	1.00 26.15	Α	N
10	MOTA	408	CA	ALA	152		-59.442	93.836	1.00 25.48	Α	С
	MOTA	409	CB	ALA	152		-59.064	94.544	1.00 25.10	Α	С
	MOTA	410	С	ALA	152		-58.290	93.937	1.00 24.90	A	С
	MOTA	411	0	ALA	152		-58.073	94.985	1.00 23.72	A	0
45	MOTA	412	N	PRO	153		-57.544	92.841	1.00 24.54	A	N
15	MOTA	413	CD	PRO	153		-57.751	91.463	1.00 25.59	Α	С
	MOTA	414	CA	PRO	153		-56.435	92.919	1.00 24.98	Α	C
	ATOM	415	CB	PRO	153		-55.859	91.495	1.00 25.27	Α	С
	MOTA	416	CG	PRO	153		-56.374	90.870	1.00 25.95	A	С
	MOTA	417	С	PRO	153		-55.421	93.993	1.00 25.30	Α	С
20	ATOM	418	0	PRO	153		~55.182	94.220	1.00 25.69	Α	0
	MOTA	419	N	VAL	154		-54.845	94.666	1.00 24.93	Α	N
	MOTA	420	CA	VAL	154		-53.884	95.730	1.00 25.63	Α	С
	ATOM	421	СВ	VAL	154		-54.044	96.884	1.00 26.14	Α	С
05	MOTA	422		VAL	154		-53.454	96.484	1.00 26.07	A	C
25	MOTA	423		VAL	154		-53.391	98.148	1.00 27.60	Α	С
	ATOM	424	C	VAL	154		-52.432	95.245	1.00 25.24	A	С
	ATOM	425	0	VAL	154		-51.531	95.980	1.00 25.07	Α	0
	ATOM	426	N	LEU	155		-52.214	94.010	1.00 24.31	Α	N
20	ATOM	427	CA	LEU	155		-50.875	93.418	1.00 24.23	Α	С
30	ATOM	428	СВ	LEU	155		-50.977	91.910	1.00 24.21	Α	С
	ATOM	429	CG	LEU	155		-49.663	91.118	1.00 25.59	Α	C
	MOTA	430		LEU	155		-48.739	91.744	1.00 24.34	Α	С
	MOTA	431		LEU	155		-49.962	89.658	1.00 24.80	Α	С
25	MOTA	432	C	LEU	155		-50.038	93.661	1.00 23.18	Α	С
35	MOTA	433	0	LEU	155		-48.915	94.149	1.00 23.96	A	0
	MOTA	434	N	PRO	156		-50.569	93.331	1.00 22.69	A	Ŋ
	ATOM	435	CD	PRO	156		-51.848	92.684	1.00 22.19	A	С
	ATOM	436	CA	PRO	156		-49.757	93.571	1.00 22.08	A	C
40	ATOM	437	CB	PRO	156		-50.698	93.171	1.00 22.24	A	С
40	MOTA	438	CG	PRO	156		-51.539	92.086	1.00 22.14	A	С
	ATOM	439	C	PRO	156		-49.277	95.031	1.00 22.20	A	С
	ATOM	440	0	PRO	156		-48.122	95.287	1.00 20.87	A	0
	MOTA	441	N	LEU	157		-50.164	95.980	1.00 20.93	A	N
45	ATOM	442	CA	LEU	157		-49.813	97.397	1.00 20.80	A	C
45	ATOM	443	CB	LEU	157		-51.046	98.269	1.00 19.44	A	С
	ATOM	444	CG	LEU	157		-50.793	99.783	1.00 19.55	A	С
	ATOM	445		LEU	157		-50.152	100.220	1.00 16.57	A	С
	ATOM	446		LEU	157		-52.113	100.522	1.00 16.20	A	С
50	ATOM	447	C	LEU	157		-48.738	97.715	1.00 20.74	A	С
50	ATOM	448	0	LEU	157		-47.774	98.422	1.00 20.79	A	0
	MOTA	449	N	VAL	158		-48.911	97.186	1.00 20.78	A	N
	MOTA	450	CA	VAL	158		-47.962	97.387	1.00 20.77	A	С
	MOTA	451	CB	VAL	158		-48.480	96.704	1.00 21.54	A	C
55	ATOM	452		VAL	158		-47.402	96.720	1.00 21.06	A	C
55	MOTA	453		VAL	158		-49.743	97.420	1.00 20.78	Α	C
	MOTA	454	C	VAL	158		-46.583	96.810	1.00 21.47	A	С
	ATOM	455	0	VAL	158		-45.549	97.428	1.00 21.74	A	0
	ATOM	456	N	THR	159	-2.293	-46.575	95.621	1.00 21.17	Α	N

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	MOTA	457		THR	159		-45.334	94.956	1.00 21.04	A	С
	MOTA	458	CB	THR	159		-45.606	93.484	1.00 21.88	Α	С
	MOTA	459	OG1	THR	159	-2.012	-46.413	92.882	1.00 22.76	A	0
	ATOM	460	CG2	THR	159	-3.143	-44.308	92.710	1.00 21.05	Α	С
5	MOTA	461	С	THR	159	-3.841	-44.677	95.682	1.00 20.47	Α	С
	ATOM	462	0	THR	159	-3.944	-43.449	95.721	1.00 20.97	Α	0
	ATOM	463	N	HIS	160	-4.721	-45.500	96.252	1.00 19.61	Α	N
	ATOM	464	CA	HIS	160	-5.866	-45.009	97.008	1.00 20.01	Α	С
	ATOM	465	CB	HIS	160	-6.789	-46.161	97.423	1.00 20.08	A	С
10	ATOM	466	CG	HIS	160		-45.744	98.354	1.00 20.94	Α	С
	ATOM	467	CD2		160	-8.085	-45.999	99.669	1.00 20.95	A	C
	MOTA	468	ND1		160		-44.925	97.965	1.00 21.19	A	N
	ATOM	469	CE1		160		-44.691	99.000	1.00 20.93	A	C
	ATOM	470	NE2		160		-45.331		1.00 20.20	A	N
15	ATOM	471	C	HIS	160		-44.308	98.254	1.00 19.95	A	C
••	ATOM	472	0	HIS	160		-43.219	98.600	1.00 20.42	A	Ö
	MOTA	473	N	PHE	161		-44.933	98.920	1.00 20.42	A	N
		474	CA	PHE	161		-44.345		1.00 18.56		C
	MOTA		CB							A	
20	ATOM	475		PHE	161		<b>-45.327</b>		1.00 18.23	A	C
20	ATOM	476	CG	PHE	161		-46.356		1.00 17.32	A	C
	MOTA	477		PHE	161		-46.314		1.00 16.89	A	C
	MOTA	478		PHE	161		-47.362		1.00 18.24	A	C
	MOTA	479		PHE	161		-47.255		1.00 17.84	A	C
05	ATOM	480		PHE	161		-48.308		1.00 17.78	Α	С
25	ATOM	481	CZ	PHE	161		-48.253		1.00 16.70	Α	С
	MOTA	482	C	PHE	161		-43.030	99.758	1.00 19.43	Α	С
	MOTA	483	0	PHE	161		-42.041		1.00 18.14	Α	0
	MOTA	484	N	ALA	162		-43.019	98.651	1.00 19.79	Α	N
	MOTA	485	CA	ALA	162		-41.809	98.233	1.00 21.41	Α	С
30	MOTA	486	CB	ALA	162		-42.061	96.923	1.00 20.88	Α	С
	MOTA	487	С	ALA	162	-2.598	-40.674	98.040	1.00 22.02	Α	С
	MOTA	488	0	ALA	162		~39.545	98.471	1.00 22.29	A	0
	ATOM	489	N	ASP	163	-3.718	-40.991	97.399	1.00 22.30	A	N
	ATOM	490	CA	ASP	163	-4.766	-40.012	97.144	1.00 22.87	Α	С
35	ATOM	491	CB	ASP	163	-5.841	-40.626	96.236	1.00 23.81	Α	С
	ATOM	492	CG	ASP	163	-5.360	40.804	94.788	1.00 25.70	Α	C
	ATOM	493	OD1	ASP	163	-5.966	-41.605	94.044	1.00 24.91	A	0
	ATOM	494	OD2	ASP	163	-4.380	-40.134	94.390	1.00 25.54	Α	0
	ATOM	495	С	ASP	163	-5.399	-39.468	98.426	1.00 23.27	Α	С
40	ATOM	496	0	ASP	163	-5.426	-38.254	98.633	1.00 22.56	Α	0
	ATOM	497	N	ILE	164	-5.896	-40.346		1.00 22.58	Α	N
	ATOM	498	CA	ILE	164			100.527	1.00 22.00	Α	
	ATOM	499	СВ	ILE	164			101.261	1.00 21.65	Α	C
	ATOM	500		ILE	164			100.302	1.00 19.34	Α	C
45	ATOM	501		ILE	164			101.807	1.00 20.52	A	Ċ
	ATOM	502		ILE	164			102.530	1.00 18.21	A	Č
	ATOM	503	C	ILE	164			101.490	1.00 22.19	A	Č
	ATOM	504	Ö	ILE	164			102.281	1.00 21.74	A	ŏ
	ATOM	505	N	ASN	165			101.425	1.00 21.74	A	N
50	ATOM	506	CA	ASN	165			102.289	1.00 22.10	A	C
50	ATOM	507	CB	ASN	165			102.200	1.00 22.10	A	
		508	CG		165			102.200	1.00 20.87		C
	ATOM			ASN						A	С
	MOTA	509 510		ASN	165			103.784	1.00 19.71	A	0
EE	ATOM	510		ASN	165			102.703	1.00 19.72	A	N
55	ATOM	511	С	ASN	165			101.858	1.00 22.73	A	С
	MOTA	512	0	ASN	165			102.690	1.00 22.46	A	0
	MOTA	513	N	THR	166			100.549	1.00 23.63	A	N
	MOTA	514	CA	THR	166	-2.758	-35.994	99.975	1.00 24.53	A	С

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	MOTA	515		THR	166		-36.082	98.448	1.00 25.05	Α	С
	MOTA	516		THR	166		-36.830	98.166	1.00 25.95	Α	0
	MOTA	517		THR	166		-34.689	97.834	1.00 24.40	Α	С
_	MOTA	518		THR	166		-35.168		1.00 24.71	A	C
5	MOTA	519		THR	166		-34.020		1.00 25.33	A	0
	ATOM	520		PHE	167		-35.765		1.00 23.87	Α	N
	MOTA	521		PHE	167		-35.125		1.00 23.69	Α	С
	MOTA	522		PHE	167		-36.132		1.00 23.28	Α	С
	MOTA	523	CG	PHE	167		-35.699		1.00 23.25	Α	С
10	MOTA	524	CD1	PHE	167		-34.629		1.00 23.28	Α	С
	MOTA	525	CD2	PHE	167		-36.351		1.00 22.82	Α	С
	MOTA	526	CE1	PHE	167	-10.800	-34.213	100.757	1.00 21.97	Α	С
	MOTA	527	CE2	PHE	167			102.427	1.00 22.68	A	C
	MOTA	528	CZ	PHE	<b>167</b> .	-11.298	-34.872	101.880	1.00 22.73	A	C
15	ATOM	529	С	PHE	167			101.755	1.00 24.07	Α	С
	ATOM	530	0	PHE	167			101.996	1.00 24.74	Α	0
	MOTA	531	N	MET	168	-6.161	-35.508	102.702	1.00 22.88	A	N
	ATOM	532	CA	MET	168	-6.133	-35.168	104.123	1.00 22.54	Α	С
	ATOM	533	CB	MET	168	-5.844	-36.420	104.974	1.00 20.50	Α	С
20	MOTA	534	CG	MET	168	-7.020	-37.378	105.090	1.00 19.43	Α	С
	MOTA	535	SD	MET	168	-6.792	-38.627	106.404	1.00 16.80	A	S
	MOTA	536	CE	MET	168	-5.899	-39.880	105.499	1.00 19.32	Α	С
	MOTA	537	С	MET	168	-5.137	-34.065	104.472	1.00 22.25	Α	C
	MOTA	538	0	MET	168	-5.459	-33.160	105.226	1.00 21.58	A	0
25	ATOM	539	N	VAL	169	-3.928	-34.144	103.932	1.00 22.92	A	N
	MOTA	540	CA	VAL	169			104.186	1.00 23.69	A	С
	MOTA	541	СВ	VAL	169			103.383	1.00 24.08	Α	С
	ATOM	542		VAL	169			103.555	1.00 25.04	Α	С
	MOTA	543	CG2	VAL	169			103.863	1.00 24.66	A	C
30	ATOM	544	С	VAL	169			103.794	1.00 24.06	Α	C
	MOTA	545	0	VAL	169	-3.335	-30.759	104.546	1.00 23.08	A	0
	MOTA	546	N	LEU	170	-4.084	-31.656	102.616	1.00 24.09	A	N
	ATOM	547	CA	LEU	170			102.131	1.00 24.47	Α	С
	ATOM	548	СВ	LEU	170			100.702	1.00 25.76	A	C
35	ATOM	549	CG	LEU	170		-30.969		1.00 27.02	A	С
	ATOM	550		LEU	170		-31.173		1.00 27.49	A	С
	ATOM	551	CD2	LEU	170		-29.881		1.00 27.84	Α	C
	ATOM	552	С	LEU	170	-5.772	-29.919	103.054	1.00 24.53	Α	C
	ATOM	553	0	LEU	170			103.246	1.00 24.40	Α	0
40	ATOM	554	N	GLN	171	-6.519	-30.855	103.637	1.00 23.66	Α	N
	ATOM	555	CA	GLN	171			104.550	1.00 23.20	Α	С
	MOTA	556	СВ	GLN	171			104.856	1.00 22.89	A	
	ATOM	557	CG	GLN	171			103.657	1.00 22.15	Α	С
	ATOM	558	CD	GLN	171			103.108	1.00 21.39	Α	C
45	ATOM	559		GLN	171			103.777	1.00 20.79	Α	Ō
. •	ATOM	560		GLN	171			101.888	1.00 20.38	A	N
	ATOM	561	C	GLN	171			105.841	1.00 23.71	Α	C
	ATOM	562	Ō	GLN	171			. 106.485	1.00 22.77	A	ō
	ATOM	563	Ŋ	VAL	172			106.232	1.00 24.17	A	N
50	ATOM	564	CA	VAL	172			107.445	1.00 25.12	A	C
	MOTA	565	CB	VAL	172			107.893	1.00 25.29	A	C
	ATOM	566		VAL	172			109.034	1.00 23.34	A	C
	ATOM	567		VAL	172			108.339	1.00 23.34	A	C
	MOTA	568	C	VAL	172			107.193	1.00 25.52	A	
55	MOTA	569	Ö	VAL	172			108.057	1.00 25.32	A	
00	MOTA	570	N	ILE	173			106.037		A	
	ATOM	571	CA	ILE	173			105.646		A	
	ATOM	572	CB	ILE	173			2 104.231		A	
	AIOH	312	CD	-111	1,7	3.002	-20.50	. TO4.431	1.00 20.70	~	C

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	MOTA	573	CG2	ILE	173		-25.491		1.00 26.79	A	С
	MOTA	574	CG1	ILE	173		-27.822		1.00 27.16	Α	C
	MOTA	575	CD1	ILE	173		-27.937		1.00 27.08	Α	С
_	ATOM	576	С	ILE	173		-25.919		1.00 27.47	Α	C
5	ATOM	577	0	ILE	173		-24.847		1.00 27.14	Α	0
	ATOM	578	N	LYS	174	-5.962	-26.305	105.122	1.00 27.73	Α	N
	MOTA	579	CA	LYS	174		-25.473		1.00 28.31	Α	С
	ATOM	580	CB	LYS	174		-26.123		1.00 28.44	Α	С
_	ATOM	581	CG	LYS	174	-7.935	-26.108	102.803	1.00 29.80	Α	С
10	ATOM	582	CD	LYS	174	-8.878	-26.964	101.975	1.00 31.27	Α	С
	ATOM	583	CE	LYS	174	-10.279	-26.412	101.958	1.00 32.16	Α	С
	MOTA	584	NZ	LYS	174	-11.147	-27.278	101.116	1.00 34.46	Α	N
	ATOM	585	С	LYS	174		-25.232		1.00 28.11	Α	С
	MOTA	586	0	LYS	174	-8.142	-24.147	106.885	1.00 28.60	Α	0
15	MOTA	587	N	PHE	175	-7.499	-26.240	107.429	1.00 27.63	Α	N
	MOTA	588	CA	PHE	175	-7.898	-26.142	108.834	1.00 27.68	Α	С
	ATOM	589	СВ	PHE	175	-7.736	-27.507	109.524	1.00 27.35	Α	С
	MOTA	590	CG	PHE	175	-7.842	-27.452	111.029	1.00 27.54	Α	С
	ATOM	591	CD1	PHE	175	-9.041	-27.119	111.651	1.00 27.14	Α	С
20	ATOM	592	CD2	PHE	175	-6.730	-27.709	111.822	1.00 27.58	Α	С
	ATOM	593	CE1	PHE	175			113.039	1.00 26.74	Α	C
	ATOM	594	CE2	PHE	175			113.215	1.00 27.74	A	C
	ATOM	595	CZ	PHE	175			113.823	1.00 27.47	A	C
	ATOM	596	C	PHE	175			109.551	1.00 28.52	A	c
25	MOTA	597	O	PHE	175			110.268	1.00 27.62	A	ō
	ATOM	598	N	THR	176			109.343	1.00 29.79	A	N
	ATOM	599	CA	THR	176			109.985	1.00 31.23	A	C
	ATOM	600	CB	THR	176			109.789	1.00 30.67	Α	c
	ATOM	601	OG1		176			108.399	1.00 30.30	A	ō
30	ATOM	602	CG2		176			110.306	1.00 30.30	A	Ċ
00	ATOM	603	C	THR	176			109.479	1.00 32.33	A	C
	MOTA	604	Ö	THR	176			110.232	1.00 32.53	A	0
	MOTA	605	N	LYS	177			108.207	1.00 32.01	A	N
	ATOM	606	CA	LYS	177			107.602	1.00 33.30	A	C
35	ATOM	607	CB	LYS	177			106.105	1.00 34.34	A	c
33	ATOM	608	CG	LYS	177			105.310	1.00 33.34		C
		609	CD	LYS	177			105.310	1.00 38.15	A A	C
	ATOM		CE	LYS	177		-21.393				
	MOTA	610			177			104.434	1.00 40.78 1.00 41.40	A	C
40	ATOM	611	NZ	LYS						A	N
40	ATOM	612	C	LYS	177			108.243	1.00 34.59	A	C
	MOTA	613	0	LYS	177			108.153	1.00 35.04	A	0
	ATOM	614	N	ASP	178			108.874	1.00 34.14		N
	ATOM	615	CA	ASP	178			109.543	1.00 34.02	A	C
AE	MOTA	616	CB	ASP	178			109.445	1.00 34.24	A	C
45	MOTA	617	CG	ASP	178			108.129	1.00 34.92	A	C
	MOTA	618		ASP	178			107.277	1.00 35.28	Α	0
	MOTA	619		ASP	178			107.950	1.00 36.67	Α	0
	MOTA	620	C	ASP	178			111.013	1.00 33.84	Α	С
	MOTA	621	0	ASP	178			111.752	1.00 34.10	Α	0
50	ATOM	622	N	LEU	179			111.439	1.00 33.64	Α	Ŋ
	ATOM	623	CA	LEU	179			2 112.819	1.00 33.48	Α	С
	ATOM	624	СВ	LEU	179			113.398	1.00 32.20	A	С
	MOTA	625	CG	LEU	179			7 113.240	1.00 31.98	Α	С
	ATOM	626		LEU	179			113.963	1.00 31.56	A	С
55	ATOM	627	CD2	LEU	179	-8.107	-22.876	113.793	1.00 30.73	Α	
	MOTA	628	С	LEU	179			112.834	1.00 33.94	Α	
	MOTA	629	0	LEU	179	-4.913	-18.982	2 112.464	1.00 33.94	A	
	ATOM	630	N	PRO	180			3 113.252	1.00 34.34	Α	

	ATOM	631	CD	PRO	180		-17.743		1.00 34		A	С
	ATOM	632	CA	PRO	180		-16.564		1.00 34		Α	С
	MOTA	633	CB	PRO	180		-15.725		1.00 34	. 81	Α	C
_	MOTA	634	CG	PRO	180		-16.250		1.00 34		Α	С
5	ATOM	635	С	PRO	180		-16.606		1.00 35	.01	A	С
	ATOM	636	0	PRO	180	-3.680	-16.112	113.320	1.00 34	.47	Α	0
	MOTA	637	N	VAL	181	-4.501	-17.208	115.092	1.00 35	.81	Α	N
	MOTA	638	CA	VAL	181	-3.200	-17.275	115.745	1.00 36		Α	С
	ATOM	639	CB	VAL	181		-17.889		1.00 37		A	C
10	ATOM	640	CG1	VAL	181		-18.014		1.00 38		A	C
	MOTA	641		VAL	181		-17.007		1.00 38		A	C
	MOTA	642	C	VAL	181		-18.051		1.00 36		A	C
	ATOM	643	ō	VAL	181		-17.730		1.00 36		A	õ
	ATOM	644	N	PHE	182		-19.069		1.00 37		Α	N
15	ATOM	645	CA	PHE	182		-19.847		1.00 37		A	C
. •	ATOM	646	СВ	PHE	182		-21.141		1.00 37		A	C
	ATOM	647	CG	PHE	182		-21.141					
	ATOM	648		PHE	182				1.00 37		A	C
							-22.806		1.00 37		A	C
20	MOTA	649		PHE	182		-21.805		1.00 37		A	С
20	MOTA	650		PHE	182			111.663	1.00 36		Α	С
	MOTA	651	CE2		182		-22.511		1.00 37		Α	С
	MOTA	652	CZ	PHE	182		-23.368		1.00 36		Α	С
	ATOM	653	С	PHE	182		-19.036		1.00 38		Α	С
0.5	ATOM	654	0	PHE	182			111.826	1.00 37		Α	0
25	MOTA	655	N	ARG	183			111.499	1.00 39	.10	Α	N
	MOTA	656	CA	ARG	183	-1.948	-17.630	110.314	1.00 40	.37	Α	C
	MOTA	657	CB	ARG	183	-3.268	-17.242	109.661	1.00 40	.60	Α	C
	ATOM '	658	CG	ARG	183	-3.094	-16.612	108.314	1.00 41	.01	Α	С
	ATOM	659	CD	ARG	183	-4.196	-17.041	107.394	1.00 40	.77	A	С
30	MOTA	660	NE	ARG	183	-4.070	-16.387	106.101	1.00 40	.69	Α	N
	ATOM	661	CZ	ARG	183	-5.001	-16.419	105.160	1.00 39		Α	С
	ATOM	662	NH1	ARG	183			105.370	1.00 39		Α	N
	ATOM	663	NH2		183			104.017	1.00 38		A	N
	ATOM	664	С	ARG	183			110.618	1.00 41		A	C
35	ATOM	665	0	ARG	183			109.747	1.00 41		A	ō
	ATOM	666	N	SER	184			111.858	1.00 42		Α	N
	ATOM	667	CA	SER	184			112.292	1.00 43		Α	C
	ATOM	668	СВ	SER	184			113.620	1.00 44		A	C
	ATOM	669	OG	SER	184			113.959	1.00 47		A	o
40	ATOM	670	c	SER	184			112.439	1.00 44		A	C
	ATOM	671	Ö	SER	184			112.538	1.00 44		A	0
	ATOM	672	N	LEU	185			112.459	1.00 44			
	ATOM	673	CA	LEU	185		-	112.578			A	
	ATOM	674	CB	LEU	185			112.945	1.00 44 1.00 43		A	C
45	ATOM	675	CG	LEU	185			114.282			A	C
43									1.00 44		A	С
	ATOM	676		LEU	185			114.362	1.00 42		A	С
	MOTA	677		LEU	185			115.423	1.00 43		A	C
	ATOM	678	C	LEU	185			111.248	1.00 45		Α	С
<b>5</b> 0	MOTA	679	0	LEU	185			110.194	1.00 45		A	0
50	ATOM	680	N	PRO	186			111.278	1.00 46		Α	N
	ATOM	681	CD	PRO	186			112.418	1.00 46		A	C
	MOTA	682	CA	PRO	186			110.018	1.00 46		Α	C
	MOTA	683	CB	PRO	186			110.477	1.00 46		A	C
	MOTA	684	CG	PRO	186			111.741	1.00 47		Α	С
55	MOTA	685	C	PRO	186	5.390	-17.348	109.144	1.00 47		Α	С
	MOTA	686	0	PRO	186			109.659	1.00 48		Α	0
	ATOM	687	N	ILE	187			107.830	1.00 47		A	N
	MOTA	688	CA	ILE	187			106.891	1.00 47		A	C
										_		_

	MOTA	689	CB	ILE	187	5.236	-17.749	105.423	1:00 48.13	A	С
	MOTA	690	CG2		187		-17.105		1.00 48.93	Α	C
	ATOM	691			187	4.975	-18.895	104.442	1.00 48.18	Α	С
	MOTA	692	CD1	ILE	187	6.169	-19.808	104.180	1.00 48.34	. A	C
5	MOTA	693	C	ILE	187	6.055	-19.460	107.095	1.00 47.81	Α	С
	MOTA	694	0	ILE	187	5.614	-20.602	106.967	1.00 47.87	Α	0
	MOTA	695	N	GLU	188	7.327	-19.221	107.405	1.00 47.20	A	N
	ATOM	696	CA	GLU	188		-20.320		1.00 46.46	A	C
	ATOM	697	СВ	GLU	188		-19.800		1.00 47.64	A	C
10	ATOM	698	CG	GLU	188		-18.469		1.00 50.01	A	C
	ATOM	699	CD	GLU	188		-17.284		1.00 51.33	A	č
	ATOM	700		GLU	188		-17.060		1.00 51.74	A	ō
	ATOM	701		GLU	188		-16.582		1.00 52.98	A	ō
	ATOM	702	C	GLU	188		-21.182		1.00 45.35	A	c
15	ATOM	703	ō	GLU	188		-22.407		1.00 44.56	A	ŏ
	MOTA	704	N	ASP	189		-20.544		1.00 44.44	A	N
	ATOM	705	CA	ASP	189		-21.271		1.00 43.61	A	C
	ATOM	706	СВ	ASP	189		-20.307		1.00 44.85	A	c
	ATOM	707	CG	ASP	189		-19.883		1.00 45.63	A	C
20	ATOM	708		ASP	189		-19.099		1.00 45.83	A	o
20	ATOM	709		ASP	189		-20.339		1.00 40.80		0
	ATOM	710	C	ASP	189		-22.070		1.00 47.10	A n	c
	MOTA	711	Ö	ASP	189		-23.142		1.00 42.00	A A	0
	ATOM	712	N	GLN	190		-21.541		1.00 42.20	A	N
25	ATOM	713	CA	GLN	190		-22.230		1.00 41.71	A	C
20	ATOM	714	СВ	GLN	190		-21.346		1.00 41.03	A	C
	MOTA	715	CG	GLN	190		-21.340		1.00 40.82	A	C
	ATOM	716	CD	GLN	190			108.348	1.00 40.62	A	C
	MOTA	717		GLN	190		-18.158		1.00 40.05	A	o
30	MOTA	718	NE2		190			107.350	1.00 41.13	A	N
00	MOTA	719	C	GLN	190		-23.514		1.00 40.60	A	C
	MOTA	720	ŏ	GLN	190			108.964	1.00 40.55	A	0
	MOTA	721	N	ILE	191			107.850	1.00 40.63	A	И
	ATOM	722	CA	ILE	191			107.095	1.00 40.39	A	C
35	ATOM	723	СВ	ILE	191			106.142	1.00 41.16	A	C
•	ATOM	724	CG2		191			105.325	1.00 41.08	A	C
	ATOM	725	CG1		191			105.216	1.00 42.55	A	C
	ATOM	726	CD1		191			104.324	1.00 43.94	A	C
	ATOM	727	C	ILE	191			108.061	1.00 39.72	A	C
40	ATOM	728	ō	ILE	191			108.009	1.00 40.14	A	ŏ
	ATOM	729	N	SER	192			108.944	1.00 38.77	A	N
	ATOM	730	CA	SER	192			109.918	1.00 38.58	Α	
	ATOM	731	СВ	SER	192			110.836	1.00 39.73	A	C
	ATOM	732	OG	SER	192			110.072	1.00 41.62	Α	ŏ
45	ATOM	733	C	SER	192			110.760	1.00 36.86	A	c
	MOTA	734	Ō	SER	192			110.868	1.00 36.31	A	ŏ
	ATOM	735	N	LEU	193			111.353	1.00 35.10	A	Ŋ
	ATOM	736	CA	LEU	193			112.181	1.00 34.68	A	C
	ATOM	737	СВ	LEU	193			112.821	1.00 34.16	Α	Č
50	ATOM	738	CG	LEU	193			113.887	1.00 34.02	A	Č
	ATOM	739		LEU	193			114.445	1.00 33.54	A	c
	ATOM	740		LEU	193			115.005	1.00 33.52	Α	C
	ATOM	741	C	LEU	193			111.395	1.00 34.25	A	C
	ATOM	742	Ö	LEU	193			111.888	1.00 32.86	A	0
55	ATOM	743	N	LEU	194			110.169	1.00 32.50	A	N
	ATOM	744	CA	LEU	194			109.317	1.00 35.38	A	C
	ATOM	745	СВ	LEU	194			108.025	1.00 36.27	A	C
	ATOM	746	CG	LEU	194			107.161	1.00 30.27	A	C
										n	•

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		747	001		104	0 300	00 000	107 000	1 00 00 00	_	_
	ATOM	747	CD1		194		-27.777		1.00 36.63	A	С
	MOTA	748		LEU	194		-27.162		1.00 39.17	Α	С
	MOTA	749	C	LEU	194	3.061	-29.420	108.979	1.00 35.03	Α	C
_	MOTA	750	0	LEU	194		-30.470		1.00 34.17	Α	0
5	ATOM	751	N	LYS	195	4.342	-29.376	108.626	1.00 34.97	Α	N
	MOTA	752	CA	LYS	195	5.079	-30.587	108.285	1.00 34.93	Α	С
	ATOM	753	СВ	LYS	195		-30.241	107.796	1.00 36.64	A	C
	ATOM	754	CG	LYS	195		-29.420		1.00 38.77	A	Ċ
	ATOM	755	CD	LYS	195			106.017	1.00 40.24	A	c
10	ATOM	756	CE	LYS	195		-28.509		1.00 40.24		C
10	MOTA	757	NZ	LYS	195		-29.167			A	
									1.00 42.42	A	N
	MOTA	758	C	LYS	195		-31.510		1.00 33.81	A	С
	MOTA	759	0	LYS	195		-32.729		1.00 33.63	Α	0
4 =	MOTA	760	N	GLY	196		-30.921		1.00 32.20	A	N
15	MOTA	761	CA	GLY	196		-31.713		1.00 30.60	Α	С
	MOTA	762	С	GLY	196		-32.292		1.00 29.37	Α	С
	MOTA	763	0	GLY	196	4.400	-33.342	113.085	1.00 28.60	Α	0
	ATOM	764	N	ALA	197	3.202	-31.648	112.210	1.00 27.66	Α	N
	ATOM	765	CA	ALA	197	1.958	-32.119	112.804	1.00 26.31	Α	C
20	MOTA	766	CB	ALA	197	1.400	-31.028	113.699	1.00 26.25	Α	С
	ATOM	767	С	ALA	197		-32.666		1.00 25.33	Α	С
	ATOM	768	0	ALA	197		-33.329		1.00 24.11	Α	ō
	ATOM	769	N	ALA	198		-32.396		1.00 24.59	Α	N
	ATOM	770	CA	ALA	198		-32.860		1.00 23.71	Α	C
25	MOTA	771	СВ	ALA	198			108.248	1.00 21.84	A	c
20	ATOM	772	C	ALA	198			109.901	1.00 21.64		C
		773	o	ALA	198			110.102		A	
	MOTA								1.00 22.27	A	0
	MOTA	774	N	VAL	199			109.825	1.00 21.45	Α	N
20	MOTA	775	CA	VAL	199			109.988	1.00 20.82	A	С
30	ATOM	776	CB	VAL	199			109.682	1.00 20.86	Α	С
	ATOM	777		VAL	199			110.025	1.00 20.51	Α	С
	ATOM	778		VAL	199			108.201	1.00 19.73	Α	С
	MOTA	779	С	VAL	199			111.387	1.00 21.29	Α	C
	ATOM	780	0	VAL	199			111.539	1.00 20.92	Α	0
35	ATOM	781	N	GLU	200	-0.012	-36.353	112.406	1.00 20.35	Α	N
	ATOM	782	CA	GLU	200	-0.463	-36.549	113.775	1.00 21.19	Α	С
	ATOM	783	CB	GLU	200	0.444	-35.780	114.747	1.00 21.68	Α	С
	ATOM	784	CG	GLU	200			114.891	1.00 22.67	Α	C
	ATOM	785	CD	GLU	200		-35.615		1.00 23.90	A	Č
40	ATOM	786	OE1		200			116.690	1.00 23.20	Α	ō
. •	ATOM	787	OE2		200			115.897	1.00 25.78	A	ŏ
	ATOM	788	C	GLU	200			113.930	1.00 23.70	A	C
	ATOM	789	Ö	GLU	200			114.466	1.00 20.46	A	
	ATOM	790	N	ILE	201			113.459	1.00 20.40		0
45		791	CA							A	N
40	ATOM			ILE	201			113.531	1.00 21.26	A	C
	ATOM	792	CB	ILE	201			112.901	1.00 21.50	Α	С
	MOTA	793		ILE	201			112.825	1.00 22.57	Α	С
	MOTA	794		ILE	201			113.735	1.00 21.48	Α	C
	MOTA	795		ILE	201			113.133	1.00 21.35	Α	C
50	MOTA	796	С	ILE	201			112.806	1.00 21.13	Α	С
	MOTA	797	0	ILE	201			113.278	1.00 21.64	Α	0
	MOTA	798	N	CYS	202	-4.136	-35.737	111.657	1.00 20.20	Α	N
	ATOM	799	CA	CYS	202			110.893	1.00 20.37	Α	С
	ATOM	800	СВ	CYS	202			109.585	1.00 20.42	A	C
55	ATOM	801	SG	CYS	202			108.393	1.00 21.08	Α	S
	MOTA	802	Ċ	CYS	202			111.694	1.00 18.85	A	C
	ATOM	803	ŏ	CYS	202			111.687	1.00 18.40	A	0
	ATOM	804	N	HIS	203			112.384	1.00 18.40		
	ALOM	004	14	1113	203	-3.401	-50.403	114.384	1.00 10.00	Α	N

	ATOM	805	CA	HIS	203	-4.781	-39.635	113.190	1.00 18.29	A	С
	MOTA	806	CB	HIS	203	-3.505	-40.312	113.686	1.00 17.63	Α	С
	MOTA	807	CG	HIS	203	-2.837	-41.157	112.646	1.00 18.10	Α	С
	ATOM	808	CD2	HIS	203	-1.678	-40.986	111.967	1.00 17.36	Α	С
5	MOTA	809	ND1	HIS	203	-3.398	-42.318	112.161	1.00 17.64	Α	N
	ATOM	810	CE1	HIS	203	-2.616	-42.825	111.225	1.00 17.59	Α	С
	MOTA	811	NE2	HIS	203	-1.567	-42.035	111.087	1.00 17.70	Α	N
	MOTA	812	С	HIS	203	-5.690	-39.255	114.361	1.00 18.57	Α	C
	MOTA	813	0	HIS	203	-6.586	-40.012	114.724	1.00 17.74	Α	0
10	ATOM	814	N	ILE	204	-5.470	-38.080	114.945	1.00 17.95	Α	N
	MOTA	815	CA	ILE	204	-6.326	-37.634	116.031	1.00 18.97	Α	С
	ATOM	816	CB	ILE	204	-5.867	-36.271	116.595	1.00 18.95	Α	C
	MOTA	817	CG2	ILE	204	-6.949	-35.696	117.530	1.00 17.63	Α	С
	MOTA	818	CG1	ILE	204	-4.529	-36.436	117.322	1.00 17.08	Α	С
15	ATOM	819	CD1	ILE	204		-35.142		1.00 16.88	Α	С
	ATOM	820	С	ILE	204		-37.491		1.00 19.25	A	С
	ATOM	821	0	ILE	204		-37.949		1.00 19.34	A	0
	ATOM	822	N	VAL	205		-36.859		1.00 19.23	A	N
	ATOM	823	CA	VAL	205			113.708	1.00 19.58	A	C
20	ATOM	824	CB	VAL	205		-35.782		1.00 19.63	A	C
	ATOM	825	CG1		205			111.680	1.00 20.05	A	c
	ATOM	826		VAL	205			112.823	1.00 19.24	A	Č
	ATOM	827	C	VAL	205			113.330	1.00 19.64	A	Č
	ATOM	828	ō	VAL	205			113.545	1.00 19.82	A	ŏ
25	ATOM	829	N	LEU	206			112.773	1.00 19.63	A	N
	ATOM	830	CA	LEU	206			112.342	1.00 19.49	A	C
	ATOM	831	СВ	LEU	206			111.379	1.00 19.58	A	c
	ATOM	832	CG	LEU	206			109.879	1.00 13.30	A	C
	ATOM	833		LEU	206			109.455	1.00 20.56	A	C
30	ATOM	834		LEU	206			109.094	1.00 20.30	A	C
00	ATOM	835	C	LEU	206			113.469	1.00 20.01		C
	ATOM	836	ŏ	LEU	206			113.242	1.00 18.04	A A	0
	ATOM	837	N	ASN	207			114.678	1.00 18.00		Ŋ
	ATOM	838	CA	ASN	207			115.817	1.00 17.72	A	C
35	ATOM	839	СВ	ASN	207			117.089	1.00 17.71	A	
00	ATOM	840	CG	ASN	207			117.009		A	C
	ATOM	841		ASN	207			119.233	1.00 17.11	A	С
		842		ASN	207			119.233	1.00 16.94	A	0
	ATOM ATOM	843	C	ASN	207			116.255	1.00 14.15	A	N
40	ATOM	844	0	ASN	207			116.032	1.00 18.08	A	C
70	ATOM	845	N	THR	207			115.782	1.00 17.24	A	0
		846	ÇA	THR					1.00 17.73	A	N
	ATOM	847	CB		208			115.978 115.973	1.00 19.30		
	ATOM			THR	208				1.00 19.32	Α	C
45	ATOM	848		THR	208			114.875	1.00 20.78		0
45	ATOM	849		THR	208			117.284	1.00 20.86		С
	MOTA	850	C	THR	208			114.987	1.00 19.41		C
	ATOM	851	0	THR	208			115.150	1.00 19.80		
	ATOM	852	N	THR	209			113.964	1.00 19.05		N
EΩ	MOTA	853	CA	THR	209			113.045	1.00 19.58		С
50	ATOM	854	CB	THR	209			111.608	1.00 19.77		С
	MOTA	855	OG1		209			111.584	1.00 19.93		0
	ATOM	856	CG2		209			111.073	1.00 20.26		С
	MOTA	857	C	THR	209			113.537	1.00 19.57		С
	MOTA	858	0	THR	209			113.079	1.00 19.72		0
55	MOTA	859	N	PHE	210			114.483	1.00 19.02		
	MOTA	860	CA	PHE	210			115.006	1.00 20.29		_
	ATOM	861	CB	PHE	210			115.873	1.00 19.30		-
	MOTA	862	CG	PHE	210	-9.938	-47.354	116.129	1.00 19.82	Α	С

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	ATOM	863	CD1	PHE	210	-9.215	-47.948	115.096	1.00 19.46	A	С
	MOTA	864	CD2	PHE	210	-9.891	-47.926	117.399	1.00 18.74	A	С
	MOTA	865	CE1	PHE	210	-8.454	-49.096	115.325	1.00 19.60	Α	С
	MOTA	866	CE2	PHE	210	-9.135	-49.072	117.641	1.00 19.21	Α	C
5	MOTA	867	CZ	PHE	210		-49.661		1.00 19.22	Α	С
	MOTA	868	С	PHE	210	-13.035	-47.176	115.810	1.00 21.23	A	С
	MOTA	869	0	PHE	210	-13.639	-46.667	116.756	1.00 20.44	Α	0
	MOTA	870	N	CYS	211	-13.229	-48.430	115.421	1.00 22.24	Α	N
	MOTA	871	CA	CYS	211	-14.175	-49.304	116.087	1.00 24.03	Α	C
10	ATOM	872	CB	CYS	211	-14.950	-50.117	115.049	1.00 24.67	Α	С
	MOTA	873	SG	CYS	211	-16.182	-51.252	115.747	1.00 25.21	Α	S
	ATOM	874	С	CYS	211	-13.385	-50.228	117.006	1.00 24.78	Α	С
	ATOM	875	0	CYS	211	-12.508	-50.962	116.562	1.00 23.56	Α	0
	ATOM	876	N	LEU	212	-13.690	-50.179	118.294	1.00 25.72	Α	N
15	MOTA	877	CA	LEU	212	-12.989	-51.006	119.262	1.00 27.59	Α	С
	ATOM	878	СВ	LEU	212	-13.380	-50.578	120.671	1.00 27.74	Α	С
	ATOM	879	CG	LEU	212	-12.881	-49.185	121.058	1.00 28.59	Α	С
	MOTA	880	CD1	LEU	212	~13.486	-48.759	122.396	1.00 28.93	Α	С
	ATOM	881		LEU	212	-11.365			1.00 27.26	Α	С
20	ATOM	882	С	LEU	212		-52.489		1.00 29.09	Α	С
	ATOM	883	0	LEU	212	-12.374	-53.322	119.189	1.00 28.72	Α	0
	ATOM	884	N	GLN	213			118.765	1.00 30.49	Α	N
	ATOM	885	CA	GLN	213			118.583	1.00 32.47	Α	С
	ATOM	886	СВ	GLN	213			118.283	1.00 35.49	Α	С
25	MOTA	887	CG	GLN	213		-55.523	118.581	1.00 40.38	A	С
	ATOM	888	CD	GLN	213			120.073	1.00 42.74	Α	C
	ATOM	889	OE1		213			120.806	1.00 44.72	Α	Ō
	ATOM	890	NE2		213			120.528	1.00 44.13	A	N
	ATOM	891	C	GLN	213			117.474	1.00 31.71	A	C
30	ATOM	892	Ö	GLN	213			117.662	1.00 31.33	A	ō
00	ATOM	893	N	THR	214			116.319	1.00 30.92	Α	N
	ATOM	894	CA	THR	214			115.183	1.00 30.21	A	C
	ATOM	895	СВ	THR	214			113.898	1.00 30.65	A	Č
	ATOM	896	OG1		214			113.707	1.00 30.08	A	ō
35	ATOM	897	CG2		214			113.983	1.00 30.64	A	Ċ
00	ATOM	898	C	THR	214			114.922	1.00 29.58	A	c
	MOTA	899	Ö	THR	214			114.079	1.00 29.04	A	ŏ
	MOTA	900	N	GLN	215			115.625	1.00 29.13	A	N
	MOTA	901	CA	GLN	215			115.454	1.00 29.85	A	c
40	ATOM	902	СВ	GLN	215			115.775	1.00 31.30	A	č
40	ATOM	903	CG	GLN	215			116.476	1.00 35.10	A	C
		904	CD		215			117.919	1.00 36.36	A	Č
	ATOM ATOM	905		GLN	215			118.547	1.00 36.52	A	ō
	ATOM	906		GLN	215			118.456	1.00 37.92	A	N
45	ATOM	907		GLN	215			114.007	1.00 29.28	A	C
73	MOTA	908		GLN	215			113.397	1.00 29.33	A	
	ATOM	909		ASN	216			113.37	1.00 27.92	A	
	ATOM	910		ASN	216			112.104	1.00 27.75	A	
		911		ASN	216			111.276	1.00 27.73	A	
50	MOTA	912		ASN	216			110.966	1.00 23.04	A	
JU	ATOM	912		L ASN	216			3 110.566	1.00 31.15	A	
	MOTA	913		2 ASN	216			3 111.130	1.00 32.04	A	
	ATOM	914		ASN	216			9 112.109	1.00 32.04	A	
	MOTA				216			5 112.109 5 113.079	1.00 25.04		
EE	MOTA	916		ASN						A A	_
55	MOTA	917		PHE	217 217			7 111.012 7 110.868	1.00 24.54 1.00 23.68	A	
	MOTA	918		PHE						A N	_
	ATOM	919		PHE	217			5 110.247		A	_
	MOTA	920	CG	PHE	217	-3.2/1	40./2.	3 111.132	1.00 21.44	Α	C

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	ATOM	921	CD1	PHE	217	-8.295	-47.709	111.143	1.00 20.23	Α	С
	MOTA	922	CD2	PHE	217	-9.127	-45.625	111.977	1.00 21.42	Α	С
	MOTA	923	CE1	PHE	217	-7.191	-47.603	111.982	1.00 21.65	Α	С
_	MOTA	924	CE2	PHE	217		-45.509		1.00 21.20	A	C
5	MOTA	925	CZ	PHE	217		-46.497		1.00 20.69	Α	С
	MOTA	926	С	PHE	217	-12.821	-47.677	109.944	1.00 23.24	Α	С
	MOTA	927	0	PHE	217	-12.714	-48.053	108.778	1.00 23.15	Α	0
	MOTA	928	N	LEU	218	-13.976	-47.292	110.475	1.00 22.85	Α	N
.0	MOTA	929	CA	LEU	218	-15.217			1.00 22.28	Α	С
10	MOTA	930	СВ	LEU	218	-16.388	-47.702	110.591	1.00 22.69	A	С
	ATOM	931	CG	LEU	218	-16.185	-49.017	111.344	1.00 24.13	Α	C
	ATOM	932	CD1	LEU	218	-17.413	-49.316	112.191	1.00 24.57	Α	С
	ATOM	933	CD2	LEU	218	-15.923	-50.148	110.346	1.00 24.80	Α	C
	MOTA	934	С	LEU	218	-15.478	-45.919	109.110	1.00 21.34	Α	C
15	MOTA	935	0	LEU	218	-15.830	-44.984	109.830	1.00 20.56	Α	0
	MOTA	936	N	CYS	219	-15.305	-45.805	107.798	1.00 21.05	Α	N
	MOTA	937	CA	CYS	219	-15.502	-44.541	107.101	1.00 20.80	Α	C
	MOTA	938	ÇВ	CYS	219	-14.203	-44.136	106.399	1.00 20.03	Α	С
	MOTA	939	SG	CYS	219	-12.762	-44.055	107.502	1.00 21.17	Α	S
20	ATOM	940	С	CYS	219	-16.640	-44.667	106.087	1.00 21.14	Α	С
	ATOM	941	0	CYS	219			104.889	1.00 20.85	A	0
	ATOM	942	N	GLY	220	-17.865	-44.492	106.574	1.00 21.18	Α	N
	ATOM	943	CA	GLY	220	-19.024	-44.612	105.710	1.00 21.35	Α	С
	ATOM	944	С	GLY	220			105.222	1.00 21.61	Α	С
25	ATOM	945	0	GLY	220			106.025	1.00 21.96	Α	0
	ATOM	946	N	PRO	221	-19.197	-46.270	103.908	1.00 21.46	Α	N
	ATOM	947	CD	PRO	221			102.816	1.00 21.21	A	C
	ATOM	948	CA	PRO	221			103.402	1.00 22.15	A	C
	ATOM	949	СВ	PRO	221			102.027	1.00 21.42	A	Č
30	MOTA	950	CG	PRO	221			101.576	1.00 20.84	A	C
	MOTA	951	С	PRO	221			103.320	1.00 22.19	A	C
	MOTA	952	Ó	PRO	221			102.938	1.00 22.79	A	Õ
	ATOM	953	N	LEU	222			103.678	1.00 21.79	A	N
	ATOM	954	CA	LEU	222			103.633	1.00 21.31	A	C
35	ATOM	955	СВ	LEU	222			103.093	1.00 19.09	A	c
	ATOM	956	CG	LEU	222			101.644	1.00 18.95	A	C
	ATOM	957		LEU	222			101.319	1.00 18.64	Α	Č
	ATOM	958		LEU	222			100.691	1.00 16.82	A	Ċ
	ATOM	959	C	LEU	222			104.999	1.00 21.86	A	c
40	ATOM	960	Ö	LEU	222			106.030	1.00 21.48	Α	o
. •	ATOM	961	N	ARG	223			104.979	1.00 21.57	Α	Ŋ
	ATOM	962	CA	ARG	223			106.175	1.00 22.37	A	C
	ATOM	963	СВ	ARG	223			106.465	1.00 24.98	Α	c
	MOTA	964	CG	ARG	223			107.490	1.00 28.92	A	C
45	ATOM	965	CD	ARG	223			106.858	1.00 32.82	A	c
	ATOM	966	NE	ARG	223			106.324	1.00 36.01	A	N
	MOTA	967	CZ	ARG	223			105.057	1.00 36.66	A	C
	ATOM	968		ARG	223			104.167	1.00 36.15	A	N
	ATOM	969	NH2		223			104.677	1.00 36.63	A	N
50	ATOM	970	С	ARG	223			105.967	1.00 21.18	A	C
00	ATOM	971	Õ	ARG	223			105.070	1.00 20.16	A	0
	ATOM	972	N	TYR	224			106.782	1.00 20.10	A	N
	ATOM	973	CA	TYR	224			106.782	1.00 18.92	A	
	ATOM	974	CB	TYR	224	-9.088		106.078	1.00 17.44		
55	MOTA	975	CG	TYR	224	-9.470		105.716		A A	
-	ATOM	976		TYR	224			105.804		A	
	ATOM	977		TYR	224			103.741	1.00 16.45	A	
									1.00 16.85	A	
	MOTA	978	CDZ	TYR	224	-0./01	-4/./5]	104.402	1.00 16.10	Α	С

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		070			004		46 000				
	MOTA	979	-	TYR	224		-46.897		1.00 16.65	Α	С
	MOTA	980	CZ	TYR	224	-10.225			1.00 16.90	Α	С
	MOTA	981	OH	TYR	224		-45.234		1.00 16.57	Α	0
_	MOTA	982	С	TYR	224	-9.354	-50.949	107.836	1.00 18.68	Α	C
5	MOTA	983	0	TYR	224	-9.712	-50.713	108.986	1.00 18.12	Α	0
	ATOM	984	N	THR	225		-51.951		1.00 17.86	Α	N
	MOTA	985	CA	THR	225		-52.894		1.00 18.12	A	C
	ATOM	986	СВ	THR	225		-54.355		1.00 16.87	A	Č
	ATOM	987			225		-54.637		1.00 16.37		
10	ATOM	988	CG2	THR	225		-54.578			A	0
10									1.00 15.99	A	C
	ATOM	989	C	THR	225		-52.781		1.00 17.74	Α	С
	ATOM	990	0	THR	225		-52.164		1.00 17.42	Α	0
	MOTA	991	N	ILE	226		-53.404		1.00 16.95	Α	N
4.5	MOTA	992	CA	ILE	226		-53.390		1.00 17.61	Α	С
15	MOTA	993	CB	ILE	226	-4.161	-53.985	111.246	1.00 17.39	Α	С
	MOTA	994	CG2	ILE	226		-55.476		1.00 15.65	Α	С
	MOTA	995	CG1	ILE	226	-2.685	-53.713	111.555	1.00 17.84	Α	С
	ATOM	996	CD1	ILE	226	-2.309	-54.002	112.989	1.00 18.23	Α	С
	MOTA	997	С	ILE	226	-3.823	-54.153	108.714	1.00 17.93	A	С
20	ATOM	998	0	ILE	226		-53.874		1.00 16.65	A	Ō
	ATOM	999	N	GLU	227		-55.089		1.00 17.96	Α	N
	ATOM	1000	CA	GLU	227		-55.840		1.00 19.11	A	C
	ATOM	1001	СВ	GLU	227		-56.981		1.00 10.11		C
	ATOM	1002	CG	GLU	227		-58.231			A	
25	ATOM	1002	CD	GLU	227		-57.975		1.00 20.80	A	C
25		1003		GLU					1.00 21.93	A	C
	ATOM				227		-57.220		1.00 23.31	Α	0
	MOTA	1005		GLU	227		-58.536		1.00 21.48	Α	0
	MOTA	1006	С	GLU	227		-54.914		1.00 19.73	Α	С
00	MOTA	1007	0	GLU	227		-55.147		1.00 18.85	Α	0
30	MOTA	1008	N	ASP	228		-53.877		1.00 18.55	Α	N
	MOTA	1009	CA	ASP	228	-4.371	-52.928	104.533	1.00 19.17	Α	C
	MOTA	1010	CB	ASP	228	-5.504	-51.886	104.522	1.00 19.08	Α	С
	MOTA	1011	CG	ASP	228	-6.846	-52.496	104.159	1.00 19.07	Α	С
	MOTA	1012	OD1	ASP	228	-6.873	-53.316	103.219	1.00 20.93	Α	0
35	ATOM	1013	OD2	ASP	228			104.795	1.00 18.63	A	0
	ATOM	1014	С	ASP	228			104.691	1.00 18.73	A	Ċ
	ATOM	1015	0	ASP	228			103.715	1.00 19.17	A	ŏ
	ATOM	1016	N	GLY	229			105.922	1.00 18.10	A	N
	ATOM	1017	CA	GLY	229		-51.253		1.00 17.50	A	C
40	ATOM	1018	C	GLY	229	-0.245		105.104	1.00 17.30		
	MOTA	1019	Ö	GLY	229			105.321		A	C
	MOTA	1020	N	ALA	230				1.00 15.48	A	0
								106.410	1.00 17.33	A	N
	MOTA	1021	CA	ALA	230			106.247	1.00 18.15	A	С
A E	MOTA	1022	CB	ALA	230			106.987	1.00 16.66	Α	С
45	MOTA	1023	C	ALA	230			104.770	1.00 19.00	Α	С
	ATOM	1024	0	ALA	230			104.344	1.00 19.76	Α	0
	ATOM	1025	N	ARG	231			103.990	1.00 19.49	Α	N
	MOTA	1026	CA	ARG	231			102.566	1.00 19.81	Α	С
	ATOM	1027	CB	ARG	231	-1.491	-55.499	101.973	1.00 20.78	Α	С
50	ATOM	1028	CG	ARG	231	-2.159	-56.765	102.524	1.00 22.44	Α	С
	MOTA	1029	CD	ARG	231	-1.366	-58.040	102.205	1.00 24.31		C
	ATOM	1030	NE	ARG	231			100.783	1.00 26.32	A	N
	MOTA	1031	CZ	ARG	231			100.143	1.00 28.16	A	C
	ATOM	1032		ARG	231			100.780	1.00 27.73	A	
55	ATOM	1032		ARG	231		-59.233		1.00 27.73		N
-	ATOM	1033	C	ARG	231			101.721		A	N
									1.00 19.74	A	C
	MOTA	1035	0	ARG	231			100.650	1.00 20.20	A	0
	ATOM	1036	N	VAL	232	U.608	-52.867	102.170	1.00 18.61	Α	N

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	MOTA	1037	CA	VAL	232		-51.826		1.00 17.26	Α	С
	MOTA	1038	CB	VAL	232		-50.434		1.00 17.52	Α	С
	MOTA	1039	CG1	VAL	232		-50.569		1.00 17.25	Α	C
_	MOTA	1040	CG2	VAL	232	0.563	-49.875	102.879	1.00 16.49	Α	С
5	MOTA	1041	С	VAL	232		-51.670		1.00 17.95	Α	C
	MOTA	1042	0	VAL	232	3.469	-50.800	101.358	1.00 17.18	Α	0
	MOTA	1043	N	GLY	233	3.189	-52.510	102.788	1.00 18.73	Α	N
	MOTA	1044	CA	GLY	233	4.582	-52.442	103.205	1.00 19.65	Α	С
	MOTA	1045	С	GLY	233		-51.997		1.00 20.28	Α	C
10	MOTA	1046	0	GLY	233	6.129	-52.078	104.969	1.00 20.44	Α	0
	MOTA	1047	N	PHE	234	3.999	-51.514	105.397	1.00 19.59	Α	N
	MOTA	1048	CA	PHE	234	4.313	-51.107	106.764	1.00 20.47	Α	С
	MOTA	1049	CB	PHE	234	3.134	-50.366	107.401	1.00 20.71	Α	С
	MOTA	1050	CG	PHE	234	2.949	-48.963	106.901	1.00 21.67	A	С
15	MOTA	1051		PHE	234		-48.602		1.00 21.54	Α	С
	MOTA	1052	CD2	PHE	234	3.922	-47.996	107.133	1.00 22.13	Α	С
	MOTA	1053	CE1	PHE	234	1.596	-47.291	105.794	1.00 23.35	A	С
	MOTA	1054	CE2	PHE	234	3.741	-46.681	106.703	1.00 23.24	Α	C
	MOTA	1055	CZ	PHE	234	2.572	-46.328	106.032	1.00 22.54	Α	С
20	ATOM	1056	C	PHE	234	4.648	-52.322	107.633	1.00 20.13	Α	С
	MOTA	1057	0	PHE	234	4.039	-53.388	107.503	1.00 19.24	Α	0
	MOTA	1058	N	GLN	235	5.606	-52.153	108.534	1.00 20.97	Α	N
	ATOM	1059	CA	GLN	235	5.997	-53.231	109.427	1.00 22.42	Α	С
	MOTA	1060	СВ	GLN	235	7.348	-52.917	110.064	1.00 24.45	Α	С
25	MOTA	1061	CG	GLN	235	8.493	-52.968	109.069	1.00 27.92	Α	С
	ATOM	1062	CD	GLN	235	9.841	-52.727	109.709	1.00 30.93	Α	С
	MOTA	1063	OE1	GLN	235	10.843	-53.322	109.304	1.00 33.56	Α	0
	MOTA	1064	NE2	GLN	235	9.883	-51.847	110.703	1.00 30.93	Α	N
	MOTA	1065	C	GLN	235	4.932	-53.444	110.494	1.00 21.69	Α	С
30	MOTA	1066	0	GLN	235	4.303	-52.498	110.966	1.00 20.35	Α	0
	MOTA	1067	N	VAL	236	4.729	-54.702	110.863	1.00 21.75	Α	N
	MOTA	1068	CA	VAL	236	3.725	-55.064	111.850	1.00 20.74	Α	С
	MOTA	1069	CB	VAL	236	3.679	~56.585	112.035	1.00 20.71	Α	C
	ATOM	1070	CG1	VAL	236		-56.953		1.00 19.30	Α	С
35	ATOM	1071	CG2	VAL	236	3.233	-57.231	110.730	1.00 20.83	Α	С
	ATOM	1072	С	VAL	236			113.197	1.00 21.19	Α	С
	ATOM	1073	0	VAL	236			113.785	1.00 20.57	Α	0
	MOTA	1074	N	GLU	237			113.695	1.00 21.62	Α	N
	MOTA	1075	CA	GLU	237	5.400	-53.702	114.981	1.00 23.17	Α	С
40	MOTA	1076	CB	GLU	237			115.323	1.00 25.38	Α	C
	ATOM	1077	CG	GLU	237			116.535	1.00 29.56	Α	С
	MOTA	1078	CD		237			117.002	1.00 32.34	Α	С
	MOTA	1079		GLU	237			116.146	1.00 34.48	A	0
	MOTA	1080	OE2	GLU	237	8.931	-53.400	118.227	1.00 33.42	Α	0
45	MOTA	1081	С	GLU	237	4.940	-52.241	114.945	1.00 22.30	Α	С
	MOTA	1082	0	GLU	237			115.905	1.00 21.24	Α	0
	MOTA	1083	N	PHE	238			113.836	1.00 21.77	Α	N
	MOTA	1084	CA	PHE	238	4.779	-50.169	113.667	1.00 22.10	A	С
	MOTA	1085	CB	PHE	238			112.331	1.00 21.91	Α	С
50	MOTA	1086	CG	PHE	238			111.954	1.00 22.44	A	С
	MOTA	1087	CD1	PHE	238			112.622	1.00 22.09	Α	С
	MOTA	1088	CD2	PHE	238			110.961	1.00 22.65	Α	С
	ATOM	1089	CE1	PHE	238	4.409	-45.914	112.310	1.00 21.36	Α	C
	MOTA	1090		PHE	238	3.073	-47.026	110.636	1.00 22.78	A	Ċ
55	MOTA	1091	CZ	PHE	238	3.436	-45.861	111.316	1.00 22.50	A	Č
	MOTA	1092	С	PHE	238			113.673	1.00 21.89	A	Ċ
	MOTA	1093	0	PHE	238			114.378	1.00 20.84	A	ō
	ATOM	1094	N	LEU	239			112.863	1.00 22.09	A	N
									- · · •	-	

	MOTA	1095	CA	LEU	239		-51.118		1.00 22.46	Α	C
	MOTA	1096	CB	LEU	239		-52.241		1.00 21.56	Α	С
	MOTA	1097	CG	LEU	239	0.402	-51.905	110.343	1.00 23.54	Α	С
_	MOTA	1098	CD1	LEU	239	0.860	-50.519	109.926	1.00 22.90	Α	C
5	MOTA	1099	CD2	LEU	239	0.925	~52.977	109.388	1.00 21.20	Α	С
	MOTA	1100	С	LEU	239	0.577	-51.376	114.132	1.00 22.41	Α	С
	ATOM	1101	0	LEU	239	-0.441	-50.776	114.470	1.00 21.62	Α	0
	ATOM	1102	N	GLU	240		-52.262		1.00 22.26	A	N
	ATOM	1103	CA	GLU	240		-52.563		1.00 22.89	A	C
10	ATOM	1104	CB	GLU	240		-53.690		1.00 25.34		c
10	ATOM	1105	CG	GLU	240		-55.069			A	
									1.00 29.77	A	C
	ATOM	1106	CD	GLU	240		-55.498		1.00 33.75	A	С
	ATOM	1107		GLU	240		-54.811		1.00 36.81	A	0
4.5	MOTA	1108		GLU	240		-56.527		1.00 35.46	Α	0
15	MOTA	1109	С	GLU	240		-51.327		1.00 21.32	Α	С
	MOTA	1110	0	GLU	240	-0.224	-51.081	117.906	1.00 20.62	Α	0
	MOTA	1111	N	LEU	241	1.780	-50.554	117.070	1.00 20.96	Α	N
	MOTA	1112	CA	LEU	241	1.888	-49.349	117.883	1.00 22.45	Α	С
	MOTA	1113	CB	LEU	241	3.239	-48.664	117.648	1.00 24.05	Α	С
20	ATOM	1114	CG	LEU	241		-47.321		1.00 26.05	A	c
	MOTA	1115		LEU	241		-47.531		1.00 27.21	A	C
	MOTA	1116		LEU	241			117.945	1.00 27.21	A	C
	MOTA	1117	C	LEU	241			117.511			
									1.00 21.92	A	C
25	MOTA	1118	0	LEU	241			118.381	1.00 21.86	A	0
25	MOTA	1119	Ŋ	LEU	242			116.210	1.00 20.84	Α	N
	MOTA	1120	CA	LEU	242			115.677	1.00 20.79	A	С
	MOTA	1121	CB	LEU	242			114.146	1.00 19.93	Α	С
	ATOM	1122	CG	LEU	242	-0.682	-46.031	113.392	1.00 21.77	Α	С
	ATOM	1123	CD1	LEU	242	-1.129	-46.358	111.984	1.00 19.10	Α	С
30	MOTA	1124	CD2	LEU	242	-1.729	-45.184	114.085	1.00 21.40	A	С
	ATOM	1125	С	LEU	242	-1.857	-47.667	116.161	1.00 20.42	Α	С
	ATOM	1126	0	LEU	242			116.678	1.00 18.91	A	0
	ATOM	1127	N	PHE	243			115.981	1.00 20.27	A	N
	ATOM	1128	CA	PHE	243			116.386	1.00 20.87	A	C
35	ATOM	1129	СВ	PHE	243			115.775	1.00 21.22	A	c
•	MOTA	1130	CG	PHE	243			114.309	1.00 21.22	A	c
	MOTA	1131		PHE	243			113.359			
		1131							1.00 19.68	A	C
	ATOM			PHE	243			113.893	1.00 19.34	A	C
40	ATOM	1133		PHE	243			112.022	1.00 20.59	A	С
40	MOTA	1134	CE2		243			112.558	1.00 19.75	Α	С
	MOTA	1135	CZ	PHE	243			111.621	1.00 20.16	Α	С
	MOTA	1136	С	PHE	243			117.899	1.00 21.36	A	С
	MOTA	1137	0	PHE	243			118.385	1.00 20.70	Α	0
	MOTA	1138	N	HIS	244	-2.652	-49.608	118.644	1.00 21.51	Α	N
45	MOTA	1139	CA	HIS	244	-2.747	-49.586	120.101	1.00 22.59	Α	С
	ATOM	1140	CB	HIS	244	-1.448	-50.061	120.757	1.00 24.95	A	С
	MOTA	1141	CG	HIS	244			122.245	1.00 27.50	A	Č
	ATOM	1142		HIS	244			123.252	1.00 28.29	Α	č
	MOTA	1143		HIS	244			122.846	1.00 28.64	A	N
50	MOTA	1144		HIS	244			124.157	1.00 29.14		
30		1145		HIS						A	С
	MOTA				244			124.430	1.00 29.46	A	N
	MOTA	1146	C	HIS	244			120.503	1.00 21.62	A	C
	ATOM	1147	0	HIS	244			121.394	1.00 21.60	Α	0
	MOTA	1148	N	PHE	245			119.844	1.00 20.43	A	
55	MOTA	1149	CA	PHE	245			120.106	1.00 20.28	Α	С
	MOTA	1150	CB	PHE	245	-1.700	-44.935	119.173	1.00 19.48	Α	С
	ATOM	1151	CG	PHE	245			119.121	1.00 20.46	A	
	ATOM	1152	CD1	. PHE	245			120.122	1.00 20.84	Α	
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	ATOM	1153	CD2	DUE	245	2 057	-43.041	110 100	1 00 00 70	_	_
	ATOM	1154	CE1		245				1.00 20.72	A	C
							-41.284		1.00 19.78	A	C
	ATOM	1155	CE2		245		-41.721		1.00 21.00	A	С
5	MOTA	1156	CZ	PHE	245		-40.844		1.00 20.53	A	С
5	MOTA	1157	С	PHE	245		-45.446		1.00 19.25	A	С
	MOTA	1158	0	PHE	245		-44.833		1.00 19.06	Α	0
	MOTA	1159	N	HIS	246		-45.844		1.00 19.01	Α	N
	MOTA	1160	CA	HIS	246		-45.546		1.00 19.09	Α	С
40	MOTA	1161	CB	HIS	246		-45.962		1.00 18.50	A	С
10	MOTA	1162	CG	HIS	246	-5.954	-44.898	115.944	1.00 18.48	A	C
	MOTA	1163	CD2	HIS	246		-44.805		1.00 18.21	Α	C
	MOTA	1164	ND1	HIS	246	-6.665	-43.719	115.863	1.00 18.52	Α	N
	MOTA	1165	CE1	HIS	246	-6.130	-42.945	114.935	1.00 17.66	Α	С
	MOTA	1166	NE2	HIS	246	-5.100	-43.581	114.407	1.00 18.46	Α	N
15	ATOM	1167	С	HIS	246		-46.185		1.00 18.38	A	С
	MOTA	1168	0	HIS	246		-45.550		1.00 18.13	A	ō
	MOTA	1169	N	GLY	247		-47.436		1.00 19.24	A	N
	ATOM	1170	CA	GLY	247		-48.090		1.00 19.94	A	C
	ATOM	1171	C	GLY	247		-47.385		1.00 20.61	A	C
20	ATOM	1172	ō	GLY	247		-47.103		1.00 20.87	A	o
	ATOM	1173	N	THR	248		-47.084		1.00 20.87	A	
	ATOM	1174	CA	THR	248		-46.413		1.00 19.95		И
	ATOM	1175	СВ	THR	248		-46.204		1.00 20.04	A	C
	ATOM	1176		THR	248		-47.467			A	C
25	ATOM	1177		THR	248		-45.601		1.00 21.47	A	0
20	ATOM	1178	C	THR	248		-45.058		1.00 18.96	A	C
	ATOM	1179	0	THR	248				1.00 20.66	A	C
	ATOM	1180					-44.736		1.00 20.22	A	0
			N	LEU	249		-44.270		1.00 21.23	A	N
30	ATOM	1181	CA	LEU	249		-42.953		1.00 21.39	Α	С
30	MOTA	1182	CB	LEU	249		-42.178		1.00 21.91	Α	С
	MOTA	1183	CG	LEU	249		-40.751		1.00 22.61	Α	С
	ATOM	1184		LEU	249			122.494	1.00 23.51	Α	C
	MOTA	1185		LEU	249		-40.133		1.00 22.87	Α	С
25	ATOM	1186	C	LEU	249			122.569	1.00 20.82	Α	С
35	MOTA	1187	0	LEU	249			123.152	1.00 19.80	Α	0
	ATOM	1188	N	ARG	250			121.828	1.00 21.20	Α	N
	MOTA	1189	CA	ARG	250			121.644	1.00 21.98	Α	C
	MOTA	1190	СВ	ARG	250			120.573	1.00 21.92	Α	С
40	MOTA	1191	CG	ARG	250			120.158	1.00 21.63	Α	С
40	MOTA	1192	CD	ARG	250			119.412	1.00 24.25	Α	С
	MOTA	1193	NE	ARG	250			118.368	1.00 25.98	Α	N
	MOTA	1194	CZ	ARG	250	-15.285	-44.293	118.503	1.00 27.08	Α	С
	ATOM	1195	NH1	ARG	250			119.644	1.00 27.73	Α	N
	MOTA	1196	NH2	ARG	250	-16.077	-44.637	117.495	1.00 27.60	Α	N
45	ATOM	1197	С	ARG	250	-11.532	-44.580	122.936	1.00 22.80	Α	С
	ATOM	1198	0	ARG	250			123.194	1.00 22.12	Α	Ō
	ATOM	1199	N	LYS	251			123.735	1.00 23.54	Α	N
	ATOM	1200	CA	LYS	251			124.997	1.00 24.94	A	C
	ATOM	1201	СВ	LYS	251			125.650	1.00 25.92	A	Č
50	MOTA	1202	CG	LYS	251			124.961	1.00 29.26	A	Ċ
	ATOM	1203	CD	LYS	251			125.690	1.00 30.89	Α	Č
	ATOM	1204	CE	LYS	251			124.889	1.00 33.42	A	c
	ATOM	1205	NZ	LYS	251			125.462	1.00 35.42	A	
	MOTA	1206	C	LYS	251			125.992	1.00 35.04	A	N
55	ATOM	1207	Ö	LYS	251			126.932	1.00 25.07		С
-	MOTA	1208	N	LEU	252			125.793	1.00 23.41	A	0
	ATOM	1209	CA	LEU	252			126.685	1.00 24.90	A A	N
	ATOM	1210	CB	LEU	252					A	C
	AION	1210	CD	neo	454	-3.903	-41.549	126.503	1.00 23.70	Α	С

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	MOTA	1211	CG	LEU	252		-42.172		1.00 24.00	A	С
	ATOM	1212	CD1	LEU	252		-41.176		1.00 22.17	Α	С
	MOTA	1213	CD2	LEU	252		-42.599		1.00 21.31	Α	С
_	ATOM	1214	С	LEU	252	-12.457			1.00 26.07	Α	С
5	MOTA	1215	0	LEU	252	-12.822			1.00 25.67	Α	0
	ATOM	1216	N	GLN	253	-13.165	-42.027	125.410	1.00 27.09	Α	N
	MOTA	1217	CA	GLN	253	-14.454	-41.394	125.142	1.00 28.60	Α	С
	MOTA	1218	CB	GLN	253	-15.498	-41.944	126.122	1.00 30.52	A	С
	MOTA	1219	CG	GLN	253	-15.871	-43.417	125.905	1.00 34.58	Α	С
10	MOTA	1220	CD	GLN	253	-16.676	-44.004	127.072	1.00 37.32	Α	С
	ATOM	1221	OE1	GLN	253	-17.588	-43.365	127.602	1.00 38.93	Α	0
	ATOM	1222	NE2	GLN	253	-16.341	-45.228	127.466	1.00 38.76	Α	N
	MOTA	1223	С	GLN	253	-14.396	-39.866	125.246	1.00 28.46	Α	С
	MOTA	1224	0	GLN	253	-15.168	-39.256	125.986	1.00 28.32	Α	0
15	MOTA	1225	N	LEU	254	-13.487	-39.254	124.494	1.00 27.65	Α	N
	ATOM	1226	CA	LEU	254	-13.323	-37.806	124.510	1.00 27.11	Α	С
	MOTA	1227	CB	LEU	254	-12.011	-37.410	123.821	1.00 25.42	Α	С
	ATOM	1228	CG	LEU	254	-10.698	-37.954	124.384	1.00 24.68	Α	C
	MOTA	1229	CD1	LEU	254	-9.530	-37.337	123.621	1.00 23.56	Α	С
20	MOTA	1230	CD2	LEU	254	-10.599	-37.627	125.867	1.00 23.28	Α	C
	ATOM	1231	С	LEU	254	-14.467	-37.078	123.817	1.00 27.57	Α	С
	ATOM	1232	0	LEU	254	-15.163	-37.642	122.974	1.00 26.67	Α	0
	ATOM	1233	N	GLN	255			124.176	1.00 28.10	Α	N
	ATOM	1234	CA	GLN	255			123.562	1.00 29.49	A	C
25	ATOM	1235	СВ	GLN	255			124.616	1.00 31.52	Α	C
	ATOM	1236	CG	GLN	255			125.810	1.00 35.22	A	Ċ
	ATOM	1237	CD	GLN	255			126.791	1.00 38.55	A	c
	ATOM	1238		GLN	255			127.185	1.00 39.64	A	Ō
	ATOM	1239	NE2		255			127.199	1.00 40.55	A	N
30	ATOM	1240	C	GLN	255			122.496	1.00 28.88	A	c
••	MOTA	1241	ŏ	GLN	255			122.594	1.00 28.44	A	ŏ
	ATOM	1242	N	GLU	256			121.478	1.00 28.49	A	N
	ATOM	1243	CA	GLU	256			120.405	1.00 28.54	A	C
	ATOM	1244	CB	GLU	256			119.607	1.00 30.20	Α	Č
35	MOTA	1245	CG	GLU	256			118.410	1.00 33.85	A	č
•	ATOM	1246	CD	GLU	256			117.304	1.00 35.60	Α	c
	ATOM	1247		GLU	256			117.567	1.00 37.88	A	o
	ATOM	1248	OE2		256			116.174	1.00 35.27	A	o
	ATOM	1249	C	GLU	256			120.848	1.00 27.93	A	c
40	ATOM	1250	Ö	GLU	256			120.319	1.00 27.24	A	0
10	MOTA	1251	N	PRO	257			121.813	1.00 27.24	A	N
	ATOM	1252	CD	PRO	257			122.517	1.00 28.01		C
	ATOM	1253	CA	PRO	257			122.254	1.00 27.60	A	c
	MOTA	1254	CB	PRO	257			123.371	1.00 27.00	A	c
45	ATOM	1255	CG	PRO	257			123.795	1.00 29.41	A	C
40	ATOM	1256	C	PRO	257			122.698	1.00 26.72	A	c
	MOTA	1257	0	PRO	257			7 122.436	1.00 20.72	A	0
	ATOM	1258	N	GLU	258			122.450	1.00 27.13	A	N
	ATOM	1259		GLU	258			3 123.337	1.00 24.44		
50					258			7 124.813		A	C
30	MOTA	1260		GLU					1.00 24.01	A	C
	ATOM	1261		GLU	258			126.004		A	
	ATOM	1262		GLU	258			5 126.911	1.00 26.63	A	
	MOTA	1263		1 GLU	258			3 126.397	1.00 26.48	A	
E =	ATOM	1264		2 GLU	258			3 128.143	1.00 28.74	A	-
55	MOTA	1265		GLU				5 122.637	1.00 23.76	A	
	MOTA	1266		GLU				9 122.634		A	
	MOTA	1267		TYR	259			7 121.634		A	
	ATOM	1268	CA	TYR	259	-10.108	33.92	4 120.446	1.00 22.11	Α	C

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	MOTA	1269	CB	TYR	259	-11.091			1.00 21.08	A	C
	MOTA	1270	CG	TYR	259	-11.261			1.00 21.43	A	C
	ATOM	1271	CD1		259	-10.198			1.00 21.24	Α	C
_	MOTA	1272	CE1		259	-10.365			1.00 19.51	Α	C
5	MOTA	1273	CD2	TYR	259		-36.622		1.00 20.98	Α	С
	MOTA	1274	CE2	TYR	259	-12.672	-38.003	120.157	1.00 21.07	Α	С
	MOTA	1275	CZ	TYR	259	-11.601	-38.860	119.914	1.00 19.86	Α	С
	MOTA	1276	OH	TYR	259	-11.788	-40.218	120.006	1.00 18.11	Α	0
	MOTA	1277	С	TYR	259	-9.466	-32.752	119.723	1.00 22.33	A	С
10	MOTA	1278	0	TYR	259	-8.308	-32.817	119.295	1.00 21.92	Α	0
	MOTA	1279	N	VAL	260	-10.229	-31.677	119.575	1.00 22.03	Α	N
	ATOM	1280	CA	VAL	260		-30.516		1.00 23.22	A	C
	ATOM	1281	СВ	VAL	260		-29.512		1.00 24.49	A	C
	ATOM	1282		VAL	260		-28.125		1.00 26.18	A	Ċ
15	ATOM	1283		VAL	260		-29.913		1.00 24.67	A	c
.0	ATOM	1284	C	VAL	260		-29.849		1.00 22.97	A	Č
	ATOM	1285	Ö	VAL	260		-29.416		1.00 22.37	A	Ö
	ATOM	1286	И	LEU	261		-29.766		1.00 22.33		N
					261		-29.161		1.00 23.08	A	
20	ATOM	1287	CA	LEU						A	C
20	MOTA	1288	CB	LEU	261		-28.975		1.00 24.90	A	C
	ATOM	1289	CG	LEU	261		-27.817		1.00 24.74	Α	C
	MOTA	1290		LEU	261		-27.932		1.00 25.84	A	С
	MOTA	1291		LEU	261		-26.501		1.00 24.89	Α	C
	ATOM	1292	С	LEU	261		-30.053		1.00 24.56	Α	C
25	MOTA	1293	0	LEU	261		-29.560		1.00 24.62	Α	0
	MOTA	1294	N	LEU	262		-31.367		1.00 24.08	Α	N
	MOTA	1295	CA	LEU	262	-5.393	-32.300	121.473	1.00 24.35	Α	С
	MOTA	1296	CB	LEU	262	-5.901	-33.739	121.558	1.00 25.52	Α	C
	MOTA	1297	CG	LEU	262	-4.940	-34.844	122.018	1.00 26.64	Α	С
30	MOTA	1298	CD1	LEU	262	-4.394	-34.531	123.404	1.00 26.40	Α	C
	MOTA	1299	CD2	LEU	262	-5.693	-36.172	122.044	1.00 26.95	Α	С
	MOTA	1300	С	LEU	262	-4.729	-32.037	120.112	1.00 23.84	Α	С
	ATOM	1301	Ο.	LEU	262	-3.504	-32.002	120.009	1.00 23.68	Α	0
	ATOM	1302	N	ALA	263			119.071	1.00 22.30	Α	N
35	MOTA	1303	CA	ALA	263			117.749	1.00 22.57	Α	С
• •	ATOM	1304	СВ	ALA	263			116.710	1.00 21.76	A	Ċ
	ATOM	1305	C	ALA	263			117.779	1.00 22.78	A	Ċ
	MOTA	1306	ŏ	ALA	263			117.186	1.00 22.16	A	ŏ
	ATOM	1307	N	ALA	264			118.470	1.00 22.57	Α	N
40	ATOM	1308	CA	ALA	264			118.586	1.00 24.00	A	c
.0	ATOM	1309	CB	ALA	264			119.332	1.00 23.05	A	c
	ATOM	1310	С	ALA	264			119.302	1.00 23.03	A	c
	ATOM	1311	Ö	ALA	264			118.944	1.00 24.41	A	_
		1311	N	MET	265			120.312	1.00 24.83		0
45	MOTA	1313	CA		265			120.312	1.00 25.02	A	N
73	MOTA			MET						A	C
	ATOM	1314	CB	MET	265			122.306	1.00 26.79	A	C
	ATOM	1315	CG	MET	265			123.415	1.00 27.47	A	C
	ATOM	1316	SD	MET	265			124.797	1.00 28.87	A	S
ΕO	MOTA	1317	CE	MET	265			125.669	1.00 27.94	A	С
50	MOTA	1318	C	MET	265			120.177	1.00 26.28	A	C
	MOTA	1319	0	MET	265			120.276	1.00 26.21	A	0
	ATOM	1320	N	ALA	266			119.314	1.00 25.89	Α	N
	ATOM	1321	CA	ALA	266			118.417		Α	
	MOTA	1322	CB	ALA	266			117.700	1.00 24.34	Α	C
55	MOTA	1323	С	ALA	266			117.402	1.00 26.44	Α	_
	MOTA	1324	0	ALA	266			117.060		A	0
	MOTA	1325	N	LEU	267	-0.292	-29.439	116.936	1.00 26.96	Α	
	MOTA	1326	CA	LEU	267	0.041	-28.401	115.968	1.00 29.21	A	

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	ATOM	1327	CB	LEU	267	-1.218	-27.636	115.547	1.00 28.28	A	С
	ATOM	1328	CG	LEU	267	-1.322	-27.086	114.116	1.00 28.22	A	С
	ATOM	1329	CD1	LEU	267	-2.153	-25.815	114.151	1.00 26.62	A	С
	MOTA	1330	CD2	LEU	267	0.040	-26.797	113.520	1.00 27.55	A	С
5	MOTA	1331	С	LEU	267		-27.405		1.00 30.31	A	C
	MOTA	1332	0	LEU	267		-27.195		1.00 30.73	Α	ō
	ATOM	1333	N	PHE	268		-26.795		1.00 31.98	A	N
	ATOM	1334	CA	PHE	268		-25.817		1.00 34.42	A	C
	ATOM	1335	СВ	PHE	268		-24.778		1.00 34.42	A	c
10	ATOM	1336	CG	PHE	268		-23.980		1.00 33.07		C
	ATOM	1337	CD1		268		-24.236			A	
	ATOM	1337		PHE	268				1.00 31.94	A	C
	ATOM	1339	CE1	PHE	268		-22.995 -23.526		1.00 32.71	A	C
									1.00 31.94	A	C
15	ATOM	1340	CE2	PHE	268		-22.278		1.00 32.75	Α	C
13	ATOM	1341	CZ	PHE	268		-22.545		1.00 32.01	Α	С
	ATOM	1342	С	PHE	268			119.171	1.00 35.98	Α	С
	MOTA	1343	0	PHE	268			120.393	1.00 36.42	Α	0
	MOTA	1344	N	SER	269			118.537	1.00 38.74	Α	N
00	MOTA	1345	CA	SER	269			119.244	1.00 41.31	Α	С
20	MOTA	1346	CB	SER	269	5.001	-29.096	118.743	1.00 41.54	Α	С
	MOTA	1347	OG	SER	269	3.849	-29.890	118.970	1.00 43.06	Α	0
	MOTA	1348	С	SER	269	6.026	-26.834	118.969	1.00 42.81	A	С
	MOTA	1349	0	SER	269	6.505	-26.782	117.840	1.00 43.23	Α	0
	MOTA	1350	N	PRO	270	6.564	-26.171	120.002	1.00 44.83	Α	N
25	ATOM	1351	CD	PRO	270	6.116	-26.232	121.406	1.00 44.95	A	С
	ATOM	1352	CA	PRO	270	7.758	-25.326	119.873	1.00 46.18	Α	C
	ATOM	1353	CB	PRO	270			121.259	1.00 45.92	A	C
	ATOM	1354	CG	PRO	270			122.156	1.00 45.81	A	Č
	ATOM	1355	С	PRO	270			119.452	1.00 47.67	A	C
30	ATOM	1356	o	PRO	270			118.935	1.00 48.27	A	Ö
	ATOM	1357	N	ASP	271			119.672	1.00 48.67	A	N
	ATOM	1358	CA	ASP	271			119.321	1.00 49.58	A	C
	ATOM	1359	СВ	ASP	271			120.409	1.00 50.08	A	C
	ATOM	1360	CG	ASP	271			120.621	1.00 50.89	A	C
35	ATOM	1361	OD1		271			120.767	1.00 50.89	A	0
-	ATOM	1362		ASP	271			120.767	1.00 51.40		
	ATOM	1363	C	ASP	271			117.950		A	0
	ATOM	1364	0	ASP	271			117.644	1.00 50.08	A	C
	MOTA		N	ARG	272			117.124	1.00 50.01	A	0
40	ATOM	1365	CA		272			117.124	1.00 50.50	A	N
70		1366		ARG					1.00 51.11	A	C
	ATOM	1367	CB	ARG	272			115.126	1.00 50.35	A	C
	MOTA	1368	CG	ARG	272			114.325	1.00 49.50	Α	
	MOTA	1369	CD	ARG	272			115.161	1.00 48.07	Α	С
45	ATOM	1370	NE	ARG	272			115.405	1.00 47.04	Α	N
45	MOTA	1371	CZ	ARG	272			115.873	1.00 45.93	Α	С
	MOTA	1372		ARG	272			116.154	1.00 45.21	Α	N
	ATOM	1373		ARG	272			116.062	1.00 44.85	A	N
	MOTA	1374	C	ARG				115.005	1.00 51.77	Α	С
	MOTA	1375	0	ARG	272			115.015	1.00 52.46	Α	0
50	MOTA	1376	N	PRO				114.317	1.00 52.32	Α	N
	MOTA	1377	CD	PRO				114.035	1.00 52.48	Α	С
	MOTA	1378	CA	PRO				113.552	1.00 52.55	A	C
	ATOM	1379	СВ	PRO	273	12.567	-30.337	112.876	1.00 52.49	Α	С
	MOTA	1380	CG	PRO	273	11.241	-31.023	112.706	1.00 52.55	A	C
55	ATOM	1381	С	PRO				112.550	1.00 52.78	A	Č
	MOTA	1382	0	PRO				111.628	1.00 52.86	A	ō
	ATOM	1383	N	GLY				112.742	1.00 52.71	A	N
	ATOM	1384	CA	GLY				111.831	1.00 52.86	A	C
					-·-		20.000		x.00 J2.80	~	C

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	MOTA	1385	С	GLY	274	11.921	-24.428	112.306	1.00 53.31	A	С
	MOTA	1386	0	GLY	274	11.825	-23.448	111.567	1.00 52.74	A	0
	MOTA	1387	N	VAL	275		-24.451		1.00 53.86	Α	N
_	MOTA	1388	ÇA	VAL	275		-23.294		1.00 54.72	A	С
5	MOTA	1389	CB	VAL	275		-23.653		1.00 54.41	Α	С
	MOTA	1390	CG1		275		-24.806		1.00 54.40	A	C
	MOTA	1391		VAL	275		-24.010		1.00 54.73	A	С
	ATOM	1392	С	VAL	275		-22.152		1.00 55.42	Α	C
	MOTA	1393	0	VAL	275			114.694	1.00 55.34	Α	0
10	MOTA	1394	N	THR	276		-20.928		1.00 56.25	Α	N
	MOTA	1395	CA	THR	276	11.888	-19.738	114.657	1.00 57.04	Α	С
	MOTA	1396	CB	THR	276	12.019	-18.806	113.428	1.00 57.12	Α	С
	ATOM	1397	OG1		276			113.054	1.00 57.58	Α	0
	MOTA	1398	CG2	THR	276			112.252	1.00 56.94	Α	C
15	MOTA	1399	С	THR	276	11.249	-18.958	115.810	1.00 57.44	Α	C
	MOTA	1400	0	THR	276	11.939	-18.283	116.573	1.00 57.90	Α	0
	MOTA	1401	N	GLN	277	9.928	-19.058	115.929	1.00 57.58	Α	N
	MOTA	1402	CA	GLN	277			116.987	1.00 57.56	A	С
	MOTA	1403	CB	GLN	277			116.453	1.00 57.96	Α	С
20	MOTA	1404	CG	GLN	277			116.777	1.00 59.08	Α	С
	MOTA	1405	CD	GLN	277	7.893	-15.506	115.683	1.00 59.41	Α	С
	MOTA	1406		GLN	277			115.303	1.00 60.10	Α	0
	MOTA	1407	NE2	GLN	277	6.933	-14.746	115.171	1.00 59.47	Α	N
	MOTA	1408	C	GLN	277			118.125	1.00 57.42	Α	С
25	MOTA	1409	0	GLN	277	7.836	-19.507	118.629	1.00 57.26	Α	0
	MOTA	1410	N	ARG	278			118.531	1.00 57.27	Α	N
	MOTA	1411	CA	ARG	278	9.899	-21.080	119.585	1.00 57.49	A	С
	MOTA	1412	CB	ARG	278	11.279	-21.630	119.951	1.00 58.82	A	С
	MOTA	1413	CG	ARG	278			120.705	1.00 60.52	Α	С
30	MOTA	1414	CD	ARG	278	12.581	-23.453	121.124	1.00 62.18	Α	С
	MOTA	1415	NE	ARG	278	12.567	-24.877	121.465	1.00 63.94	Α	N
	MOTA	1416	CZ	ARG	278			122.495	1.00 64.51	Α	С
	MOTA	1417	NH1	ARG	278			122.697	1.00 64.55	Α	N
	MOTA	1418	NH2		278			123.328	1.00 64.96	A	N
35	MOTA	1419	C	ARG	278			120.851	1.00 56.92	Α	С
	ATOM	1420	0	ARG	278			121.380	1.00 56.95	A	0
	ATOM	1421	N	ASP	279			121.344	1.00 56.06	A	N
	ATOM	1422	CA	ASP	279			122.551	1.00 55.37	Α	С
	MOTA	1423	СВ	ASP	279			122.990	1.00 56.68	Α	С
40	MOTA	1424	CG	ASP	279			123.419	1.00 57.63	Α	С
	MOTA	1425		ASP	279			122.547	1.00 58.84	Α	0
	MOTA	1426		ASP	279			124.629	1.00 58.07	Α	_
	ATOM	1427	C	ASP	279			122.338	1.00 54.14	Α	С
4.5	MOTA	1428	0	ASP	279			123.156	1.00 53.64	Α	0
45	MOTA	1429	N	GLU	280			121.235	1.00 52.98	A	N
	MOTA	1430	CA	GLU	280			120.918	1.00 51.96	Α	С
	MOTA	1431	CB	GLU	280			119.598	1.00 52.88	Α	С
	MOTA	1432	CG	GLU	280			119.329	1.00 54.43	Α	С
50	MOTA	1433	CD	GLU	280			117.951	1.00 55.52	Α	С
50	MOTA	1434		. GLU	280			117.570	1.00 56.15	Α	0
	MOTA	1435		GLU	280			117.252	1.00 55.94	Α	0
	MOTA	1436	C	GLU	280			120.821	1.00 50.62	Α	С
	MOTA	1437	0	GLU	280			121.470	1.00 50.58	Α	
	ATOM	1438	N	ILE	281			120.004	1.00 49.04	Α	
55	MOTA	1439	CA	ILE	281			119.815	1.00 47.41	A	
	ATOM	1440		ILE	281			118.764	1.00 46.53	A	
	ATOM	1441		ILE	281			118.655	1.00 45.22	Α	
	MOTA	1442	CG1	ILE	281	5.378	-21.282	117.408	1.00 45.56	A	С

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	MOTA	1443	CD1	ILE	281	6.138	-22.010	116.326	1.00 45.01	Α	С
	MOTA	1444	С	ILE	281	4.533	-21.886	121.131	1.00 47.29	Α	С
	MOTA	1445	0	ILE	281	3.525	-22.549	121.383	1.00 46.81	A	0
_	MOTA	1446	N	ASP	282	5.568	-21.797	121.965	1.00 46.94	Α	N
5	MOTA	1447	CA	ASP	282			123.260	1.00 46.55	A	С
	ATOM	1448	CB	ASP	282	6.946	-22.311	123.926	1.00 47.68	Α	С
	ATOM	1449	CG	ASP	282	7.098	-23.166	125.170	1.00 48.85	Α	С
	ATOM	1450	OD1	ASP	282	6.921	-24.402	125.082	1.00 49.57	Α	0
	ATOM	1451	OD2	ASP	282			126.242	1.00 49.92	Α	0
10	MOTA	1452	С	ASP	282	4.470	-21.876	124.144	1.00 45.75	Α	С
	ATOM	1453	0	ASP	282	3.847	-22.583	124.934	1.00 45.36	A	0
	MOTA	1454	N	GLN	283	4.226	-20.576	124.000	1.00 45.11	Α	N
	ATOM	1455	CA	GLN	283	3.181	-19.904	124.770	1.00 45.13	Α	С
	MOTA	1456	CB	GLN	283	3.256	-18.387	124.563	1.00 46.75	Α	С
15	MOTA	1457	CG	GLN	283	2.563	-17.534	125.637	1.00 50.09	Α	С
	MOTA	1458	CD	GLN	283	1.092	-17.879	125.849	1.00 52.42	Α	С
	ATOM	1459	OE1	GLN	283	0.754	-18.757	126.649	1.00 54.11	Α	0
	MOTA	1460	NE2	GLN	283	0.210	-17.189	125.129	1.00 53.63	Α	Ŋ
	ATOM	1461	С	GLN	283	1.821	-20.423	124.292	1.00 43.58	Α	С
20	MOTA	1462	0	GLN	283			125.099	1.00 43.49	Α	0
	ATOM	1463	N	LEU	284	1.664	-20.573	122.979	1.00 42.02	A	N
	MOTA	1464	CA	LEU	284			122.428	1.00 40.90	A	C
	MOTA	1465	CB	LEU	284			120.892	1.00 40.60	A	C
	ATOM	1466	CG	LEU	284			120.110	1.00 40.12	A	C
25	ATOM	1467		LEU	284			120.918	1.00 39.50	A	C
	MOTA	1468		LEU	284			119.789	1.00 40.64	Α	Č
	ATOM	1469	C	LEU	284			122.934	1.00 39.59	A	C
	ATOM	1470	Ō	LEU	284			123.292	1.00 39.24	A	Ō
	MOTA	1471	N	GLN	285			122.950	1.00 39.22	A	N
30	MOTA	1472	CA	GLN	285			123.433	1.00 38.77	A	C
	MOTA	1473	СВ	GLN	285			123.452	1.00 39.75	A	Ċ
	ATOM	1474	CG	GLN	285			124.049	1.00 42.43	A	c
	ATOM	1475	CD	GLN	285			123.117	1.00 43.45	A	C
	MOTA	1476	OE1		285			122.730	1.00 45.03	A	ō
35	ATOM	1477		GLN	285			122.747	1.00 44.86	Α	N
-	ATOM	1478	C	GLN	285			124.843	1.00 37.86	A	C
	MOTA	1479	Ö	GLN	285			125.138	1.00 37.02	A	o.
	ATOM	1480	N	GLU	286			125.710	1.00 37.54	A	N
	ATOM	1481	CA	GLU	286			127.085	1.00 37.97	A	C
40	ATOM	1482	СВ	GLU	286			127.884	1.00 39.37	Α	Č
	ATOM	1483	CG	GLU	286			128.993	1.00 42.37	A	c
	ATOM	1484	CD	GLU	286			129.792	1.00 44.10	A	Č
	ATOM	1485		GLU	286			130.402	1.00 45.20	A	Ö
	ATOM	1486		GLU	286			129.803	1.00 46.83	A	Ö
45	MOTA	1487	С	GLU	286			127.167	1.00 36.84	A	Č
	ATOM	1488	Ö	GLU	286			128.047	1.00 36.65	A	ŏ
	ATOM	1489	N	GLU	287			126.251	1.00 37.13	A	N
	MOTA	1490	CA	GLU	287			126.203	1.00 37.20	Α	C
	MOTA	1491	CB	GLU	287			125.128	1.00 39.14	A	Č
50	ATOM	1492	CG	GLU	287			125.181	1.00 42.57	A	Č
00	MOTA	1493	CD	GLU	287			2 124.108	1.00 44.54	A	C
	ATOM	1494		GLU	287			124.049	1.00 45.67	A	Ö
	MOTA	1495	OE2		287			2 123.330	1.00 45.07	A	o
	ATOM	1496	C	GLU	287			7 125.885	1.00 36.19	A	C
55	ATOM	1497	Ö	GLU	287			125.553	1.00 35.33	A	0
55	MOTA	1498	И	MET	288			124.907	1.00 35.38	A	
	ATOM	1499	CA	MET	288			124.507	1.00 33.38		N
		1500	CB	MET	288			9 123.313	1.00 34.70	A	
	MOTA	1200	CD	PIE I	200	-5.564	-23.00	, 143.313	1.00 35.45	A	C

	ATOM	1501	CG	MET	288	-3 064	-24.841	122 100	1 00 36 40		~
	ATOM	1502	SD	MET	288		-24.110		1.00 36.49 1.00 37.77	A	c s
	ATOM	1503	CE	MET	288		-22.370			A	
	ATOM	1504	CE	MET	288		-25.930		1.00 38.91	A	C
5									1.00 33.74	A	С
3	ATOM	1505	0	MET	288		-26.379		1.00 33.02	A	0
	ATOM	1506	N	ALA	289		-26.260		1.00 33.19	A	N
	ATOM	1507	CA	ALA	289		-27.195		1.00 32.47	Α	С
	ATOM	1508	CB	ALA	289		-27.340		1.00 31.87	Α	С
10	MOTA	1509	С	ALA	289		-26.772		1.00 32.22	Α	С
10	ATOM	1510	0	ALA	289		-27.575		1.00 31.24	Α	0
	MOTA	1511	N	LEU	290		-25.512		1.00 32.24	Α	N
	MOTA	1512	CA	LEU	290		-24.985		1.00 32.90	Α	С
	MOTA	1513	СВ	LEU	290		-23.543		1.00 34.69	Α	C
	MOTA	1514	CG	LEU	290		-23.383		1.00 35.82	A	C
15	MOTA	1515		LEU	290		-21.900		1.00 36.30	A	C
	MOTA	1516	CD2	LEU	290	-2.601	-24.111	132.314	1.00 36.19	Α	C
	MOTA	1517	С	LEU	290	-5.910	-25.050	129.761	1.00 32.17	Α	С
	MOTA	1518	0	LEU	290	-6.707	-25.393	130.628	1.00 32.35	Α	0
	ATOM	1519	N	THR	291	-6.283	-24.713	128.533	1.00 31.75	Α	N
20	ATOM	1520	CA	THR	291	-7.680	-24.761	128.124	1.00 31.71	Α	С
	ATOM	1521	CB	THR	291	-7.837	-24.298	126.662	1.00 32.01	Α	C
	MOTA	1522	OG1	THR	291			126.497	1.00 32.68	A	Ō
	ATOM	1523	CG2		291			126.291	1.00 31.69	A	C
	ATOM	1524	С	THR	291			128.252	1.00 31.72	A	Ċ
25	ATOM	1525	0	THR	291			128.760	1.00 31.36	A	ō
	MOTA	1526	N	LEU	292			127.792	1.00 31.96	A	N
	ATOM	1527	CA	LEU	292			127.864	1.00 32.82	A	C
	MOTA	1528	СВ	LEU	292			127.278	1.00 31.84	A	C
	ATOM	1529	CG	LEU	292			126.636	1.00 32.12	A	C
30	MOTA	1530	CD1	LEU	292			126.649	1.00 30.55	A	C
	MOTA	1531	CD2	LEU	292			127.367	1.00 31.13	A	Ċ
	MOTA	1532	С	LEU	292			129.318	1.00 33.29	A	Ċ
	MOTA	1533	0	LEU	292			129.643	1.00 32.92	A	o
	MOTA	1534	N	GLN	293			130.185	1.00 34.69	A	N
35	ATOM	1535	CA	GLN	293			131.603	1.00 36.33	A	C
	ATOM	1536	СВ	GLN	293			132.368	1.00 37.25	A	č
	ATOM	1537	CG	GLN	293			132.047	1.00 38.64	A	C
	ATOM	1538	CD	GLN	293			132.948	1.00 39.42	A	Ċ
	MOTA	1539		GLN	293			134.169	1.00 40.32	A	o
40	ATOM	1540	NE2		293			132.355	1.00 39.89	A	N
	ATOM	1541	С	GLN	293			132.194	1.00 37.50	A	C
	ATOM	1542	ō	GLN	293			132.852	1.00 37.40	A	0
	ATOM	1543	N	SER	294			131.969	1.00 37.40	A	N
	ATOM	1544	CA	SER	294			132.480	1.00 39.65	A	C
45	ATOM	1545	СВ	SER	294			131.994	1.00 39.69	A	
. •	ATOM	1546	OG	SER	294			132.355	1.00 40.48		C
	ATOM	1547	C	SER	294			131.996	1.00 40.48	A	0
	ATOM	1548	Ö	SER	294			132.770	1.00 40.07	A	C
	ATOM	1549	N	TYR	295			132.770		A	0
50	ATOM	1550	CA	TYR	295			130.712	1.00 40.29	A	N
50	ATOM	1551	CB	TYR	295			128.633	1.00 41.02	A	C
	ATOM	1552	CG	TYR	295			127.941	1.00 40.48	A	С
									1.00 40.03	A	C
	ATOM	1553		TYR	295			128.033	1.00 39.72	A	C
55	ATOM	1554 1555	CD2	TYR TYR	295 295			127.483	1.00 40.15	A	C
55	ATOM ATOM	1556	CE2		295 295			127.271	1.00 39.70	A	C
			CEZ	TYR	295			126.722	1.00 39.35	A	C
	ATOM	1557						126.834	1.00 40.12	A	C
	ATOM	1558	OH	TYR	295	-10./48	-30.399	126.308	1.00 41.32	Α	0

	ATOM	1559	С	TYR	295	-12.507	-29 465	130.824	1.00 41.79	A	С
	ATOM	1560	Ö	TYR	295	-13.649			1.00 42.03	A	ō
	ATOM	1561	N	ILE	296	-11.455			1.00 42.99	A	N
	ATOM	1562	CA	ILE	296	-11.591			1.00 44.15	A	c
5	ATOM	1563	CB	ILE	296	-10.250			1.00 43.14	A	c
•	MOTA	1564	CG2		296	-10.230			1.00 42.45	A	c
	MOTA	1565	CG1	ILE	296			130.279	1.00 42.43	A	C
	ATOM	1566	CD1		296			130.163	1.00 42.95	A	C
	ATOM	1567	C	ILE	296			133.150	1.00 41.86	A	C
10	ATOM	1568	0	ILE	296			133.604	1.00 45.70		0
10	ATOM	1569	N	LYS	297			133.866	1.00 43.29	A	Ŋ
	ATOM	1570	CA	LYS	297			135.248	1.00 48.10	A A	C
	ATOM	1571	CB	LYS	297			135.828	1.00 51.07	A	C
	ATOM	1572	CG	LYS	297			136.112	1.00 52.35		C
15	ATOM	1573	CD	LYS	297			136.696	1.00 52.33	A.	C
10	MOTA	1574	CE	LYS	297			137.020	1.00 54.08	A	C
	ATOM	1575	NZ	LYS	297		-27.527		1.00 54.08	A	
	ATOM	1576	C	LYS	297			137.367	1.00 54.48	A	N
	MOTA	1577	0	LYS	297			136.328		A	С
20		1578			297 298				1.00 53.38	A	0
20	ATOM		N CA	GLY				134.374 134.397	1.00 55.42	A	N
	ATOM	1579		GLY	298			134.397	1.00 58.64	A	C
	MOTA	1580 1581	C	GLY	298 298				1.00 61.18	A	C
	MOTA		0	GLY				133.409 133.895	1.00 61.26	A	0
25	MOTA	1582 1583	N	GLN GLN	299				1.00 63.92	A	N
25	MOTA		CA		299			133.405	1.00 66.98	A	C
	MOTA	1584	CB	GLN	299			133.853	1.00 67.13	A	C
	ATOM	1585	CG	GLN	299			135.333	1.00 67.29	A	C
	ATOM	1586	CD	GLN	299			136.282	1.00 67.70	A	C
30	ATOM	1587		GLN	299			136.230	1.00 67.82	Α	0
30	ATOM	1588	NE2		299			137.161	1.00 67.75	A	N
	ATOM	1589	C	GLN	299			133.850	1.00 68.93	A	C
	ATOM	1590	0	GLN	299			134.831	1.00 69.45	A	0
	ATOM	1591	N	GLN	300			133.116	1.00 71.11	A	N
25	ATOM	1592	CA	GLN	300			133.446	1.00 73.14	A	C
35	ATOM	1593	CB	GLN	300			132.250	1.00 73.51	A	С
	ATOM	1594	CG	GLN	300			131.267	1.00 74.36	A	C
	ATOM	1595	CD	GLN	300			131.902	1.00 75.03	A	С
	ATOM	1596		GLN	300			132.842	1.00 75.24	A	0
40	ATOM	1597	NE2		300			131.392	1.00 75.25	A	N
40	MOTA	1598	C	GLN	300			134.661	1.00 74.34	A	C
	ATOM	1599	0	GLN	300			135.700	1.00 74.59	A	0
	MOTA	1600	N	ARG	301			134.543	1.00 75.65	A	N
	MOTA	1601	CA	ARG	301			135.667	1.00 76.68	A	C
45	MOTA	1602	CB	ARG	301			135.677	1.00 77.20	A	C
40	MOTA	1603	CG	ARG	301			137.063	1.00 78.13	A	С
	ATOM	1604	CD	ARG	301			137.011	1.00 78.94	A	С
	ATOM	1605	NE	ARG	301			138.344	1.00 79.59	A	N
	MOTA	1606	CZ	ARG	301			138.602	1.00 79.96	A	С
<b>5</b> 0	MOTA	1607		ARG	301			139.854	1.00 80.06	A	N
50	MOTA	1608		ARG	301			137.617	1.00 79.96	A	N
	ATOM	1609	C	ARG	301			135.678	1.00 76.98	A	С
	MOTA	1610	0	ARG	301			134.788	1.00 76.77	A	0
	ATOM	1611	N	ARG	302			136.713	1.00 77.23	A	N
E E	ATOM	1612	CA	ARG	302			136.920	1.00 77.33	A	С
55	ATOM	1613	CB	ARG	302			138.232	1.00 77.94	A	С
	ATOM	1614	CG	ARG	302			139.425	1.00 79.00	A	С
	ATOM	1615	CD	ARG	302			140.647	1.00 80.06	A	C
	MOTA	1616	NE	ARG	302	-19.619	-39.045	141.727	1.00 80.81	Α	Ŋ

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		_									_
	MOTA	1617		ARG	302	-20.120		_	1.00 81.28	Α	С
	MOTA	1618	NH1		302	-20.414			1.00 81.53	A	N
	MOTA	1619		ARG	302	-20.329			1.00 81.32	Α	N
_	MOTA	1620	С	ARG	302	-18.432		_	1.00 76.86	Α	С
5	ATOM	1621	0	ARG	302	-19.245			1.00 77.13	Α	0
	MOTA	1622	N	PRO	303	-17.641			1.00 76.19	Α	N
	MOTA	1623	CD	PRO	303	-17.651			1.00 76.11	Α	С
	MOTA	1624	CA	PRO	303	-16.638	-41.999	136.747	1.00 75.35	Α	С
	MOTA	1625	СВ	PRO	303	-16.150			1.00 75.81	A	С
10	MOTA	1626	CG	PRO	303	-16.298	-43.254	134.770	1.00 75.99	A	С
	MOTA	1627	С	PRO	303	-15.502	-40.983	136.888	1.00 74.26	Α	С
	MOTA	1628	0	PRO	303			135.901	1.00 74.49	A	0
	MOTA	1629	N	ARG	304	-15.251	-40.562	138.125	1.00 72.62	Α	N
	MOTA	1630	CA	ARG	304			138.422	1.00 70.74	Α	С
15	ATOM	1631	CB	ARG	304	-13.927	-39.545	139.932	1.00 71.66	Α	C
	MOTA	1632	CG	ARG	304	-13.223	-40.787	140.498	1.00 72.36	Α	С
	MOTA	1633	CD	ARG	304	-14.164	-41.984	140.613	1.00 73.27	A	C
	MOTA	1634	NE	ARG	304	-13.466	-43.229	140.936	1.00 73.97	Α	N
	MOTA	1635	CZ	ARG	304	-12.741	-43.940	140.075	1.00 74.21	Α	С
20	MOTA	1636	NH1	ARG	304	-12.601	-43.543	138.817	1.00 74.26	A	N
	MOTA	1637	NH2	ARG	304	-12.154	-45.060	140.473	1.00 74.59	Α	N
	ATOM	1638	С	ARG	304	-12.909	-40.014	137.716	1.00 68.87	A	С
	ATOM	1639	0	ARG	304	-12.746	-41.179	137.351	1.00 68.86	Α	0
	MOTA	1640	N	ASP	305	-11.995	-39.066	137.526	1.00 66.37	Α	N
25	MOTA	1641	CA	ASP	305	-10.727	-39.378	136.875	1.00 62.94	Α	C
	ATOM	1642	СВ	ASP	305	-10.978	-40.055	135.525	1.00 63.29	Α	С
	ATOM	1643	CG	ASP	305	-9.704	-40.566	134.887	1.00 63.63	Α	С
	ATOM	1644	OD1	ASP	305	-8.944	-41.283	135.570	1.00 63.64	Α	0
	MOTA	1645	OD2	ASP	305	-9.465	-40.259	133.701	1.00 64.48	Α	0
30	MOTA	1646	С	ASP	305	-9.804	-38.183	136.670	1.00 60.10	Α	С
	ATOM	1647	0	ASP	305	-9.951	-37.428	135.704	1.00 59.69	A	0
	ATOM	1648	N	ARG	306	-8.855	-38.005	137.584	1.00 56.47	A	N
	ATOM	1649	CA	ARG	306	-7.896	-36.918	137.437	1.00 52.72	Α	С
	ATOM	1650	CB	ARG	306	-7.376	-36.424	138.792	1.00 54.11	Α	С
35	ATOM	1651	CG	ARG	306	-6.551	-35.137	138.659	1.00 55.62	Α	C
	ATOM	1652	CD	ARG	306	-5.264	-35.152	139.480	1.00 57.19	Α	С
	ATOM	1653	NE	ARG	306	-5.384	-34.423	140.743	1.00 58.88	Α	N
	MOTA	1654	CZ	ARG	306	-4.358	-34.121	141.537	1.00 59.27	Α	С
	ATOM	1655	NH1	ARG	306	-3.125	-34.487	141.204	1.00 59.61	Α	N
40	ATOM	1656	NH2	ARG	306	-4.561	-33.443	142.662	1.00 59.29	A	N
	ATOM	1657	С	ARG	306	-6.721	-37.437	136.612	1.00 49.02	Α	С
	MOTA	1658	0	ARG	306	-5.719	-36.750	136.448	1.00 49.06	Α	0
	MOTA	1659	N	PHE	307	-6.857	-38.659	136.101	1.00 44.87	Α	N
	MOTA	1660	CA	PHE	307	-5.828	-39.299	135.288	1.00 40.64	Α	С
45	MOTA	1661	CB	PHE	307			135.477	1.00 39.73	Α	С
	MOTA	1662	CG	PHE	307	-5.603	-41.275	136.880	1.00 39.02	Α	C
	MOTA	1663		PHE	307			3 137.349	1.00 38.54	Α	С
	MOTA	1664		PHE	307			5 137.738	1.00 38.66	Α	C
	ATOM	1665	CE1		307			1 138.650	1.00 38.12	Α	C
50	MOTA	1666	CE2		307			2 139.044	1.00 39.04	Α	
	ATOM	1667	CZ	PHE	307			2 139.499	1.00 38.17	A	
	ATOM	1668	С	PHE	307	-5.976	-38.982	2 133.796	1.00 38.25	Α	С
	ATOM	1669		PHE	307			133.026		A	
	ATOM	1670	N	LEU	308			133.391	1.00 35.76	A	
55	ATOM	1671	CA	LEU	308			2 131.984		A	
	ATOM	1672	СВ	LEU	308			2 131.792		A	
	ATOM	1673	CG	LEU	308			B 130.351			
	ATOM	1674		LEU	308			4 129.613			
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	MOTA	1675	CD2	LEU	308	-9.036	-38.913	129.623	1.00 31.34	Α	С
	MOTA	1676	С	LEU	308	-6.389	-37.268	131.337	1.00 31.73	A	С
	MOTA	1677	0	LEU	308	-5.809	-37.601	130.306	1.00 30.35	Α	0
	MOTA	1678	N	TYR	309				1.00 29.96	A	N
5	ATOM	1679	CA	TYR	309		-35.149		1.00 28.55	A	С
	MOTA	1680	CB	TYR	309		-33.865		1.00 28.94	A	C
	ATOM	1681	CG	TYR	309		-32.828		1.00 29.27	A	C
	MOTA	1682	CD1	TYR	309		-32.250		1.00 29.24	A	C
	ATOM	1683	CE1		309		-31.315		1.00 29.07	A	Ċ
10	ATOM	1684	CD2		309		-32.443		1.00 29.31	Α	Č
	ATOM	1685	CE2	TYR	309		-31.511		1.00 29.09	A	Č
	ATOM	1686	CZ	TYR	309		-30.955		1.00 28.97	A	C
	ATOM	1687	ОН	TYR	309		-30.049		1.00 29.80	Α	ŏ
	ATOM	1688	C	TYR	309		-35.738		1.00 27.07	A	Č
15	ATOM	1689	Ö	TYR	309		-35.558		1.00 26.88	A	ŏ
. •	MOTA	1690	N	ALA	310		-36.436		1.00 25.89	Α	N
	ATOM	1691	CA	ALA	310		-37.051		1.00 25.16	A	C
	MOTA	1692	СВ	ALA	310		-37.690		1.00 24.32	A	c
	ATOM	1693	C	ALA	310		-38.096		1.00 24.96	A	C
20	ATOM	1694	ō	ALA	310		-38.142		1.00 24.12	A	0
	ATOM	1695	N	LYS	311		-38.930		1.00 24.12	A	N
	ATOM	1696	CA	LYS	311		-39.938		1.00 24.13	A	C
	ATOM	1697	CB	LYS	311		-40.752		1.00 25.00	A	C
	ATOM	1698	CG	LYS	311		-41.760		1.00 23.00		C
25	ATOM	1699	CD	LYS	311		-42.472		1.00 27.78	A A	C
20	ATOM	1700	CE	LYS	311			132.285	1.00 29.79		C
	ATOM	1701	NZ	LYS	311		-43.836		1.00 31.20	A	N
	ATOM	1702	C	LYS	311			128.508	1.00 33.30	A	С
	ATOM	1702	0	LYS	311			127.583		A	0
30	ATOM	1703	N	LEU	312			128.378	1.00 23.05	A	
30	MOTA	1704	CA	LEU	312				1.00 22.94	A	N
	ATOM	1705	CB	LEU	312			127.113 127.143	1.00 23.55	A	C
	ATOM	1707	CG	LEU	312				1.00 24.36	A	C
		1707		LEU	312			127.073	1.00 25.69	A	C
35	MOTA	1708		LEU	312			126.879	1.00 24.81	A	C
33	ATOM		CDZ					125.910 126.748	1.00 26.15	A	C
	MOTA	1710	0	LEU	312				1.00 23.28	A	C
	MOTA	1711 1712	N	LEU	312 313			125.580 127.733	1.00 22.28	A	0
	ATOM			LEU				127.733	1.00 22.16	A	N
40	MOTA	1713	CA		313				1.00 22.07	A	C
40	ATOM ATOM	1714	CB	LEU	313			128.694		A	C
		1715	CG	LEU LEU	313			129.064	1.00 21.22	A	C
	MOTA MOTA	1716 1717		LEU	313 313			130.284	1.00 20.26	A	C
								127.886	1.00 18.10	A	C
45	ATOM	1718	C	LEU	313			127.059	1.00 22.48	A	С
45	ATOM	1719	0	LEU	313			126.152	1.00 22.20	A	0
	MOTA	1720	N	GLY	314			127.733	1.00 22.89	A	N
	ATOM	1721	CA	GLY	314			127.402	1.00 23.30	A	C
	ATOM	1722	C	GLY	314			125.949	1.00 23.94	A	С
EΛ	ATOM	1723	0	GLY	314			125.200	1.00 24.54	A	0
50	ATOM	1724	N	LEU	315			125.549	1.00 23.59	A	N
	MOTA	1725	CA	LEU	315			124.182	1.00 22.89	A	С
	MOTA	1726	CB	LEU	315			124.070	1.00 23.44	Α	С
	ATOM	1727	CG	LEU	315			124.681	1.00 23.97	Α	С
EE	ATOM	1728		LEU	315			124.742	1.00 23.92	Α	С
55	ATOM	1729		LEU	315			123.854	1.00 23.72	A	C
	ATOM	1730	C	LEU	315			123.167	1.00 22.47	A	С
	MOTA	1731	0	LEU	315			122.044	1.00 22.11	Α	0
	MOTA	1732	N	LEU	316	-0.072	-37.849	123.544	1.00 22.35	Α	N

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	ATOM	1733	CA	LEU	316	0.524	-36.882	122.626	1.00 22.93	Α	С
	ATOM	1734	CB	LEU	316	0.454	-35.465	123.194	1.00 23.82	Α	С
	ATOM	1735	CG	LEU	316	-0.934	-34.817	123.164	1.00 25.69	A	C
	MOTA	1736	CD1	LEU	316	-0.879	-33.440	123.802	1.00 25.97	A	C
5	ATOM	1737	CD2	LEU	316	-1.418	-34.720	121.722	1.00 26.87	Α	C
	MOTA	1738	С	LEU	316		-37.273		1.00 23.05	A	Č
	MOTA	1739	0	LEU	316		-37.162		1.00 23.13	A	ō
	ATOM	1740	N	ALA	317		-37.741		1.00 22.68	A	N
	ATOM	1741	CA	ALA	317		-38.169		1.00 23.73	A	C
10	ATOM	1742	СВ	ALA	317		-38.439		1.00 23.73	A	C
	ATOM	1743	C	ALA	317		-39.426		1.00 23.13		
	ATOM	1744	Ö	ALA	317		-39.567		1.00 23.75	A	C
	ATOM	1745	N	GLU	318		-40.334		1.00 22.15	A	0
	ATOM	1746	CA	GLU	318		-41.562			A	N
15	ATOM	1747	CB				-42.543		1.00 26.08	A	C
15				GLU	318				1.00 28.21	Α	C
	MOTA	1748	CG	GLU	318		-43.831		1.00 32.51	A	C
	MOTA	1749	CD	GLU	318		-45.004		1.00 35.59	A	С
	MOTA	1750		GLU	318		-45.114		1.00 36.82	A	0
20	ATOM	1751	OE2		318		-45.811		1.00 38.53	A	0
20	MOTA	1752	C	GLU	318		-41.300		1.00 25.72	A	С
	MOTA	1753	0	GLU	318		-41.929		1.00 24.88	Α	0
	MOTA	1754	N	LEU	319		-40.375		1.00 25.55	A	N
	MOTA	1755	CA	LEU	319		-40.001		1.00 25.73	Α	С
0.5	MOTA	1756	CB	LEU	319		-38.994		1.00 24.34	Α	С
25	MOTA	1757	CG	LEU	319			117.311	1.00 24.57	Α	С
	MOTA	1758		LEU	319			116.426	1.00 21.78	A	С
	MOTA	1759	CD2	LEU	319	-1.131	-37.713	117.576	1.00 23.45	Α	C
	MOTA	1760	С	LEU	319	3.112	-39.395	118.103	1.00 26.16	Α	С
	MOTA	1761	0	LEU	319	3.389	-39.500	116.907	1.00 26.27	Α	0
30	MOTA	1762	N	ARG	320	3.889	-38.757	118.969	1.00 26.63	Α	N
	MOTA	1763	CA	ARG	320	5.159	-38.169	118.587	1.00 27.47	Α	С
	MOTA	1764	CB	ARG	320	5.750	-37.430	119.786	1.00 29.62	A	С
	MOTA	1765	CG	ARG	320	6.936	-36.574	119.463	1.00 32.70	Α	С
	ATOM	1766	CD	ARG	320	6.505	-35.319	118.741	1.00 35.19	Α	С
35	MOTA	1767	NE	ARG	320	7.665	-34.554	118.307	1.00 36.76	A	N
	MOTA	1768	CZ	ARG	320	7.899	-33.287	118.624	1.00 37.53	Α	С
	ATOM	1769	NH1	ARG	320	7.054	-32.609	119.389	1.00 37.83	Α	N
	MOTA	1770	NH2	ARG	320	8.993	-32.697	118.166	1.00 39.80	Α	N
	MOTA	1771	С	ARG	320			118.169	1.00 26.95	A	C
40	ATOM	1772	0	ARG	320			117.209	1.00 27.09	Α	Ō
	ATOM	1773	N	SER	321			118.898	1.00 25.51	A	N
	MOTA	1774	CA	SER	321			118.603	1.00 25.26	A	
	ATOM	1775	СВ	SER	321			119.661	1.00 24.40	A	Č
	MOTA	1776	OG	SER	321			120.843	1.00 26.98	A	Ö
45	ATOM	1777	C	SER	321			117.261	1.00 24.22	A	Č
	ATOM	1778	Ö	SER	321			116.450	1.00 23.84	A	o
	ATOM	1779	N	ILE	322			117.051	1.00 23.23	A	N
	ATOM	1780	CA	ILE	322			115.810	1.00 23.25		
	ATOM	1781	СВ	ILE	322			115.876	1.00 23.30	A	C
50	ATOM	1782		ILE	322			114.484	1.00 23.28	A	
00	ATOM	1783		ILE	322			114.464		A	C
	MOTA	1784	CD1		322			117.140	1.00 22.75	A	C
			C						1.00 21.11	A	C
	MOTA	1785		ILE	322			114.639	1.00 24.10	A	C
55	MOTA	1786	0	ILE	322			113.590	1.00 23.74	A	0
J	MOTA	1787	N	ASN	323			114.826	1.00 24.50	Α	N
	MOTA	1788	CA	ASN	323			113.787	1.00 25.76	A	С
	MOTA	1789	CB	ASN	323	5.4/7 E 000	-38.280	114.295	1.00 27.72	Α	С
	MOTA	1790	CG	ASN	323	5.909	-37.271	113.244	1.00 30.34	Α	С

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	ATOM	1791	OD1	ASN	323	6.870	-37.489	112.509	1.00 33.18	Α	0
	ATOM	1792	ND2		323		-36.148		1.00 33.10	A	N
	ATOM	1793	С	ASN	323		-40.100		1.00 25.95	A	C
	ATOM	1794	ō	ASN	323		-40.282		1.00 25.34		
5	ATOM	1795	N	GLU	324		-40.232			A	0
•	ATOM	1796	CA	GLU	324		-40.232		1.00 26.65	A	N
	ATOM	1797	CB						1.00 27.58	Α	С
		_		GLU	324		-40.535		1.00 30.21	A	С
	ATOM	1798	CG	GLU	324		-39.180		1.00 34.60	Α	С
40	ATOM	1799	CD	GLU	324		-39.163		1.00 37.84	Α	C
10	ATOM	1800		GLU	324		-40.138		1.00 39.66	Α	0
	MOTA	1801	OE2		324		-38.173		1.00 39.51	Α	0
	MOTA	1802	С	GLU	324	9.343	-41.988	113.549	1.00 26.74	Α	С
	ATOM	1803	0	GLU	324	10.238	-42.221	112.740	1.00 26.19	Α	0
	MOTA	1804	N	ALA	325	8.464	-42.914	113.923	1.00 25.09	Α	N
15	MOTA	1805	CA	ALA	325	8.517	-44.258	113.359	1.00 25.00	Α	С
	MOTA	1806	CB	ALA	325	7.619	-45.209	114.153	1.00 24.73	Α	С
	MOTA	1807	С	ALA	325		-44.238		1.00 24.89	A	C
	ATOM	1808	0	ALA	325		-45.073		1.00 25.47	A	ō
	ATOM	1809	N	TYR	326		-43.297		1.00 23.96	A	N
20	MOTA	1810	CA	TYR	326		-43.145		1.00 23.70	A	C
	ATOM	1811	СВ	TYR	326		-41.988		1.00 23.70		
	ATOM	1812	CG	TYR	326		-42.402		1.00 22.03	A	C
	MOTA	1813		TYR	326		-43.180		<del>-</del>	A	C
	MOTA	1814	CE1						1.00 20.69	Α	C
25		1815			326		-43.541		1.00 19.55	A	С
25	ATOM			TYR	326		-41.996		1.00 21.09	A	C
	ATOM	1816		TYR	326		-42.348		1.00 20.21	A	С
	ATOM	1817	CZ	TYR	326		-43.117		1.00 20.21	Α	С
	MOTA	1818	OH	TYR	326			109.749	1.00 19.33	Α	0
20	ATOM	1819	С	TYR	326			109.280	1.00 24.16	Α	С
30	MOTA	1820	0	TYR	326	8.276	-43.360	108.196	1.00 23.33	Α	0
	MOTA	1821	N	GLY	327			109.802	1.00 25.27	Α	N
	MOTA	1822	CA	GLY	327			109.116	1.00 26.47	Α	С
	MOTA	1823	С	GLY	327	11.054	-42.765	108.912	1.00 27.58	Α	C
	ATOM	1824	0	GLY	327	11.633	-42.937	107.848	1.00 27.44	Α	0
35	MOTA	1825	N	TYR	328	11.176	-43.601	109.938	1.00 29.10	Α	N
	MOTA	1826	CA	TYR	328			109.838	1.00 30.30	A	C
	MOTA	1827	СВ	TYR	328			111.182	1.00 31.95	A	Ċ
	ATOM	1828	CG	TYR	328			111.127	1.00 33.79	A	c
	ATOM	1829	CD1		328			111.361	1.00 35.24	A	C
40	ATOM	1830		TYR	328			111.242	1.00 36.06	A	c
	ATOM	1831		TYR	328			110.778	1.00 34.19	A	C
	ATOM	1832		TYR	328			110.778	1.00 34.19		
	ATOM	1833	CZ	TYR	328			110.832			C
	ATOM	1834	OH	TYR	328			110.884	1.00 36.45	A	C
45	ATOM	1835							1.00 37.83	A	0
73			C	TYR	328			108.758	1.00 30.82	A	С
	ATOM	1836	0	TYR	328			107.966	1.00 30.88	A	0
	MOTA	1837	N	GLN	329			108.737	1.00 31.03	A	N
	MOTA	1838	CA	GLN	329			107.759	1.00 31.18	A	С
50	MOTA	1839	CB	GLN	329			108.011	1.00 29.87	Α	С
50	MOTA	1840	CG	GLN	329			109.377	1.00 30.38	Α	С
	MOTA	1841	CD	GLN	329			109.523	1.00 31.26	A	С
	MOTA	1842		GLN	329			110.338	1.00 30.58	Α	0
	MOTA	1843	NE2	GLN	329			108.729	1.00 30.24	Α	N
	MOTA	1844	C	GLN	329			106.338	1.00 31.84	A	C
55	MOTA	1845	0	GLN	329			105.457	1.00 31.67	A	ō
	MOTA	1846	N	ILE	330			106.118	1.00 32.78	A	N
	ATOM	1847	CA	ILE	330			104.800	1.00 34.51	A	C
	MOTA	1848	СВ	ILE	330			104.798	1.00 34.57		
				~~~	330	2.443	-42.0JJ	104./70	1.00 34.5/	Α	С

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	MOTA	1849	CG2		330		-42.113		1.00 34.87	A	С
	MOTA	1850	CG1		330		-42.697		1.00 34.64	Α	С
	MOTA	1851		ILE	330				1.00 35.62	Α	С
_	MOTA	1852	С	ILE	330		-44.519		1.00 35.74	Α	С
5	MOTA	1853	0	ILE	330		-44.599		1.00 34.91	Α	0
	ATOM	1854	N	GLN	331		-44.604		1.00 37.46	Α	N
	MOTA	1855	CA	GLN	331		-44.776		1.00 39.69	A	C
	MOTA	1856	CB	GLN	331		-44.235		1.00 42.13	A	С
4.0	MOTA	1857	CG	GLN	331		-42.735		1.00 44.67	Α	С
10	MOTA	1858	CD	GLN	331		-42.325		1.00 47.11	Α	С
	MOTA	1859	OE1		331		-42.870		1.00 48.26	Α	0
	MOTA	1860	NE2		331		-41.362		1.00 48.66	Α	N
	MOTA	1861	С	GLN	331		-46.211		1.00 39.80	Α	С
4 =	MOTA	1862	0	GLN	331		-46.463		1.00 40.44	A	0
15	MOTA	1863	N	HIS	332		-47.155		1.00 39.86	Α	N
	ATOM	1864	CA	HIS	332		-48.546		1.00 40.67	Α	С
	MOTA	1865	CB	HIS	332		-49.161		1.00 42.81	Α	С
	MOTA	1866	CG	HIS	332		-48.524		1.00 46.28	Α	С
	MOTA	1867		HIS	332			108.086	1.00 47.01	Α	С
20	MOTA	1868	ND1		332	16.466	-48.992	107.275	1.00 47.57	Α	N
	MOTA	1869		HIS	332			107.961	1.00 47.73	Α	С
	MOTA	1870		HIS	332			108.461	1.00 48.10	Α	N
	MOTA	1871	С	HIS	332			104.337	1.00 39.76	Α	С
	MOTA	1872	0	HIS	332	13.241	-50.587	104.082	1.00 39.61	Α	0
25	MOTA	1873	N	ILE	333			103.882	1.00 38.23	A	N
	MOTA	1874	CA	ILE	333			103.055	1.00 37.05	Α	C
	MOTA	1875	СВ	ILE	333			103.779	1.00 36.60	Α	С
	MOTA	1876	CG2	ILE	333			102.878	1.00 36.04	Α	С
	ATOM	1877	CG1	ILE	333			105.066	1.00 37.04	A	С
30	ATOM	1878		ILE	333			105.963	1.00 36.04	Α	C
	ATOM	1879	С	ILE	333			101.713	1.00 36.44	A	C
	MOTA	1880	0	ILE	333			101.599	1.00 36.09	Α	0
	MOTA	1881	N	GLN	334			100.707	1.00 35.60	A	N
	MOTA	1882	CA	GLN	334		-48.909		1.00 35.11	Α	C
35	MOTA	1883	СВ	GLN	334		-49.788		1.00 37.05	Α	С
	MOTA	1884	CG	GLN	334		-49.113		1.00 40.94	Α	С
	MOTA	1885	CD	GLN	334		-48.947		1.00 43.49	Α	С
	MOTA	1886		GLN	334		-48.031		1.00 45.60	Α	0
40	MOTA	1887	NE2		334		-49.846		1.00 44.60	Α	N
40	MOTA	1888	С	GLN	334		-48.925		1.00 33.48	Α	C
	ATOM	1889	0	GLN	334		-49.929		1.00 32.53	Α	0
	MOTA	1890	N	GLY	335		-47.801		1.00 32.61	A	
	MOTA	1891	CA	GLY	335		-47.724		1.00 31.27	Α	С
45	ATOM	1892	C	GLY	335		-47.059		1.00 30.52	A	C
45	MOTA	1893	0	GLY	335		-46.519		1.00 29.45	Α	0
	ATOM	1894	N	LEU	336			100.122	1.00 30.35	Α	N
	MOTA	1895	CA	LEU	336			101.073	1.00 30.47	Α	С
	MOTA	1896	CB	LEU	336			102.515	1.00 30.60	Α	С
50	ATOM	1897	CG	LEU	336			103.265	1.00 31.59	Α	C
50	ATOM	1898		LEU	336			104.731	1.00 32.05	Α	C
	ATOM	1899		LEU	336			103.153	1.00 31.01	Α	С
	MOTA	1900	C	LEU	336			100.845	1.00 30.50	Α	С
	MOTA	1901	0	LEU	336			100.891	1.00 29.66	Α	0
	MOTA	1902	N	SER	337			100.603	1.00 31.26	Α	N
55	MOTA	1903	CA	SER	337			100.395	1.00 32.62	Α	-
	MOTA	1904	CB	SER	337			100.141	1.00 33.56	Α	
	MOTA	1905	OG	SER	337		-42.777		1.00 35.00	Α	0
	ATOM	1906	С	SER	337	6.286	-42.419	99.263	1.00 32.68	Α	С

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	MOTA	1907	0	SER	337	5.792	-41.300	99.246	1.00 33.22	Α	0
	MOTA	1908	N	ALA	338		-43.321	98.324	1.00 33.04	A	N
	MOTA	1909	CA	ALA	338	5.133	-43.016	97.218	1.00 33.69	Α	С
_:	MOTA	1910	CB	ALA	338		-44.199	96.257	1.00 33.20	Α	С
5	MOTA	1911	С	ALA	338	3.723	-42.669	97.711	1.00 34.44	Α	С
	MOTA	1912	0	ALA	338	2.979	-41.952	97.032	1.00 34.62	Α	0
	MOTA	1913	N	MET	339	3.348	-43.186	98.879	1.00 34.28	Α	N
	MOTA	1914	CA	MET	339	2.024	-42.906	99.422	1.00 35.53	Α	C
	MOTA	1915	CB	MET	339			100.400	1.00 31.99	Α	С
10	MOTA	1916	CG	MET	339		-45.292	99.701	1.00 29.56	Α	С
	ATOM	1917	SD	MET	339		-46.628		1.00 26.76	Α	S
	ATOM	1918	CE	MET	339	2.486	-47.154	101.313	1.00 25.89	Α	С
	MOTA	1919	С	MET	339	1.956	-41.537	100.086	1.00 38.11	Α	С
	MOTA	1920	0	MET	339	0.883	-41.074	100.475	1.00 38.51	A	0
15	MOTA	1921	N	MET	340	3.108	-40.891	100.217	1.00 41.19	Α	N
	MOTA	1922	CA	MET	340	3.162	-39.564	100.792	1.00 45.17	Α	С
	MOTA	1923	CB	MET	340	3.933	-39.576	102.108	1.00 45.06	Α	С
	MOTA	1924	CG	MET	340	3.761	-38.294	102.892	1.00 45.91	Α	С
	MOTA	1925	SD	MET	340			103.173	1.00 46.05	Α	S
20	ATOM	1926	CE	MET	340	1.556	-37.032	101.643	1.00 45.26	Α	С
	MOTA	1927	С	MET	340	3.848	-38.636	99.793	1.00 48.30	Α	С
	MOTA	1928	0	MET	340	4.965	-38.182	100.023	1.00 48.09	Α	0
	MOTA	1929	N	PRO	341	3.180	-38.348	98.662	1.00 51.89	A	N
	MOTA	1930	CD	PRO	341	1.818	-38.785	98.307	1.00 52.61	Α	С
25	ATOM	1931	CA	PRO	341		-37.474	97.616	1.00 55.17	Α	С
	MOTA	1932	CB	PRO	341		-37.321	96.649	1.00 54.46	Α	С
	MOTA	1933	CG	PRO	341	1.812	-38.613	96.810	1.00 53.80	Α	С
	MOTA	1934	С	PRO	341		-36.129	98.151	1.00 58.43	Α	С
	MOTA	1935	0	PRO	341		-35.838	98.177	1.00 58.94	Α	0
30	MOTA	1936	N	LEU	342	3.244	-35.312	98.573	1.00 61.89	Α	N
	ATOM	1937	CA	LEU	342	3.538	-33.993	99.108	1.00 65.33	Α	C
	MOTA	1938	CB	LEU	342	2.297	-33.094	99.015	1.00 65.46	A	С
	MOTA	1939	CG	LEU	342	1.863	-32.488	97.675	1.00 65.98	A	С
	MOTA	1940		LEU	342		-33.558	96.613	1.00 66.50	A	C
35	ATOM	1941		LEU	342		-31.702	97.907	1.00 66.63	Α	C
	MOTA	1942	С	LEU	342		-34.121		1.00 67.54	Α	С
	MOTA	1943	0	LEU	342	3.808	-35.183	101.167	1.00 67.71	Α	0
	MOTA	1944	N	LEU	343	4.490		-	1.00 70.31	Α	N
4.0	MOTA	1945	CA	LEU	343	4.909		102.505	1.00 73.10	Α	С
40	MOTA	1946	CB	LEU	343			103.401	1.00 73.06	Α	С
	MOTA	1947	CG	LEU	343			104.783	1.00 73.13	Α	С
	MOTA	1948		LEU	343			105.615	1.00 73.35	Α	C
	MOTA	1949		LEU	343			104.613	1.00 73.19	Α	С
	MOTA	1950	С	LEU	343			102.919	1.00 75.17	Α	C
45	MOTA	1951	0	LEU	343			103.242	1.00 75.75	Α	0
	MOTA	1952	N	GLN	344			102.908	1.00 77.19	Α	N
	MOTA	1953	CA	GLN	344			103.303	1.00 79.02	Α	C
	MOTA	1954	СВ	GLN	344			102.552	1.00 79.26	A	С
	MOTA	1955	CG	GLN	344			101.062	1.00 79.55	Α	С
50	MOTA	1956	CD	GLN	344			100.495	1.00 79.92	Α	С
	MOTA	1957		GLN	344		-38.044		1.00 80.00	A	0
	MOTA	1958	NE2	GLN	344			101.366	1.00 79.76	Α	N
	MOTA	1959	C	GLN	344			103.034	1.00 80.26	A	С
	MOTA	1960	0	GLN	344			102.587	1.00 80.47	A	
55	MOTA	1961	N	GLU	345			103.314	1.00 81.48	A	
	MOTA	1962	CA	GLU	345			103.093	1.00 82.57	Α	_
	MOTA	1963	CB	GLU	345			104.182	1.00 83.18	Α	С
	MOTA	1964	CG	GLU	345	12.229	-35.802	105.598	1.00 84.08	A	

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	MOTA	1965	CD	GLU	345	13.093 -36.500 106.634 1.00 84.62 A	C
	MOTA	1966	OE1		345	13.032 -37.746 106.722 1.00 85.08 A	0
	MOTA	1967		GLU	345	13.834 -35.801 107.359 1.00 84.92 A	0
_	ATOM	1968	С	GLU	345	12.106 -36.112 101.721 1.00 82.85 A	С
5	MOTA	1969	0	GLU	345	13.062 -36.916 101.657 1.00 83.15 A	0
	MOTA	1970	TXO	GLU	345	11.457 -35.728 100.723 1.00 83.01 A	0
	TER	1971		GLU	345	A	
	MOTA	1972	CB	PRO	103	12.922 -89.522 143.199 1.00 81.05 B	С
	MOTA	1973	CG	PRO	103	13.639 -89.140 144.492 1.00 81.13 B	С
10	ATOM	1974	С	PRO	103	13.827 -89.814 140.872 1.00 80.76 B	C
	ATOM	1975	0	PRO	103	13.218 -88.817 140.479 1.00 80.92 B	ō
	ATOM	1976	N	PRO	103	15.298 -89.351 142.841 1.00 81.13 B	N
	ATOM	1977	CD	PRO	103	14.976 -88.538 144.028 1.00 81.18 B	C
	ATOM	1978	CA	PRO	103	14.080 -90.046 142.362 1.00 80.95 B	C
15	MOTA	1979	N	VAL	104		
13	ATOM	1980	CA	VAL	104		N
						14.125 -90.648 138.601 1.00 79.68 B	C
	ATOM	1981	CB	VAL	104	15.488 -90.651 137.868 1.00 79.97 B	C
	MOTA	1982		VAL	104	16.297 -89.424 138.267 1.00 80.04 B	С
00	MOTA	1983		VAL	104	16.254 -91.933 138.186 1.00 79.80 B	C
20	ATOM	1984	С	VAL	104	13.296 -91.823 138.096 1.00 78.96 B	С
	MOTA	1985	0	VAL	104	13.242 -92.872 138.740 1.00 79.04 B	0
	MOTA	1986	N	GLN	105	12.654 -91.649 136.943 1.00 77.77 B	N
	MOTA	1987	CA	GLN	105	11.830 -92.710 136.373 1.00 76.25 B	С
	MOTA	1988	CB	GLN	105	10.461 -92.159 135.952 1.00 76.99 B	С
25	MOTA	1989	CG	GLN	105	9.447 -93.249 135.604 1.00 77.64 B	C
	MOTA	1990	CD	GLN	105	8.032 -92.718 135.433 1.00 78.06 B	C
	ATOM .	1991	OE1	GLN	105	7.534 -91.960 136.268 1.00 78.05 B	Ō
	ATOM	1992	NE2		105	7.371 -93.130 134.355 1.00 77.95 B	N
	ATOM	1993	C	GLN	105	12.495 -93.407 135.185 1.00 74.64 B	C
30	ATOM	1994	Ö	GLN	105	12.485 -94.634 135.110 1.00 74.66 B	0
00	ATOM	1995	N	LEU	106	13.067 -92.618 134.273 1.00 72.54 B	N
	ATOM	1996	CA	LEU	106		
	MOTA	1997	CB	LEU	106		C
						15.259 -92.860 133.160 1.00 70.25 B	C
35	MOTA	1998	CG	LEU	106	15.813 -91.432 133.205 1.00 70.18 B	C
33	ATOM	1999		LEU	106	15.723 -90.881 134.616 1.00 70.47 B	C
	MOTA	2000		LEU	106	17.267 -91.442 132.752 1.00 69.72 B	C
	MOTA	2001	С	LEU	106	13.521 -94.596 132.761 1.00 68.42 B	C
	ATOM	2002	0	LEU	106	14.449 -95.402 132.853 1.00 68.15 B	0
4.0	MOTA	2003	N	SER	107	12.295 -94.949 132.382 1.00 66.23 B	N
40	MOTA	2004	CA	SER	107	11.955 -96.333 132.064 1.00 63.76 B	С
	MOTA	2005	CB	SER	107	10.469 -96.441 131.721 1.00 63.69 B	C
	ATOM	2006	OG	SER	107	10.152 -97.713 131.185 1.00 63.42 B	0
	ATOM	2007	С	SER	107	12.782 -96.865 130.904 1.00 62.28 B	С
	ATOM	2008	0	SER	107	13.328 -96.097 130.117 1.00 62.13 B	0
45	ATOM	2009	N	LYS	108	12.878 -98.187 130.806 1.00 60.42 B	N
	MOTA	2010	CA	LYS	108	13.633 -98.817 129.733 1.00 58.34 B	C
	MOTA	2011	СВ	LYS	108	13.706-100.328 129.957 1.00 59.09 B	Ċ
	MOTA	2012	CG	LYS	108	14.716-101.043 129.078 1.00 59.82 B	Ċ
	ATOM	2013	CD	LYS	108	16.139-100.630 129.430 1.00 61.01 B	C
50	ATOM	2014	CE	LYS	108	17.167-101.382 128.589 1.00 61.73 B	C
00	ATOM	2015	NZ	LYS	108		
			C				N
	MOTA	2016		LYS	108		C
	ATOM	2017	0	LYS	108	13.524 -97.970 127.491 1.00 56.15 B	0
55	ATOM	2018	N	GLU	109	11.654 -98.895 128.342 1.00 54.51 B	N
55	MOTA	2019	CA	GLU	109	10.846 -98.673 127.148 1.00 52.68 B	С
	ATOM	2020	СВ	GLU	109	9.464 -99.323 127.297 1.00 53.23 B	С
	ATOM	2021	CG	GLU	109	9.004 -99.560 128.730 1.00 54.59 B	С
	ATOM	2022	CD	GLU	109	9.624-100.809 129.340 1.00 55.04 B	С

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	MOTA	2023	OE1		109			128.809	1.00 55.28	В	0
	MOTA	2024		GLU	109			130.343	1.00 55.07	В	0
	MOTA	2025		GLU	109			126.819	1.00 50.98	В	С
_	MOTA	2026		GLU	109			125.662	1.00 50.38	В	0
5	MOTA	2027		GLN	110			127.832	1.00 49.17	В	N
	MOTA	2028		GLN	110			127.607	1.00 47.33	В	С
	MOTA	2029		GLN	110			128.926	1.00 47.12	В	С
	ATOM	2030		GLN	110			129.606	1.00 47.16	В	С
4.0	MOTA	2031		GLN	110			130.870	1.00 47.26	В	С
10	MOTA	2032	OE1		110			131.711	1.00 46.64	В	0
	MOTA	2033		GLN	110			131.014	1.00 47.72	В	N
	MOTA	2034	С	GLN	110			126.921	1.00 46.46	В	С
	MOTA	2035	0	GLN	110			126.018	1.00 45.75	В	0
	MOTA	2036	N	GLU	111			127.343	1.00 45.39	В	N
15	MOTA	2037	CA	GLU	111			126.714	1.00 44.94	В	С
	MOTA	2038	CB	GLU	111	15.526	-95.026	127.436	1.00 46.33	В	C
	ATOM	2039	CG	GLU	111	15.591	-94.724	128.924	1.00 48.97	В	С
	ATOM	2040	CD	GLU	111			129.479	1.00 50.99	В	С
	ATOM	2041	OE1	GLU	111			129.335	1.00 51.59	В	0
20	ATOM	2042	OE2	GLU	111	17.378	-95.840	130.049	1.00 51.77	В	0
	MOTA	2043	С	GLU	111	14.292	-94.918	125.262	1.00 43.27	В	С
	MOTA	2044	0	GLU	111	14.768	-94.227	124.362	1.00 42.84	В	0
	MOTA	2045	N	GLU	112	13735	-96.107	125.050	1.00 41.39	В	N
	MOTA	2046	CA	GLU	112	13.627	-96.696	123.724	1.00 39.73	В	С
25	ATOM	2047	СВ	GLU	112	13.040	-98.104	123.833	1.00 40.89	В	С
	MOTA	2048	CG	GLU	112	12.809	-98.801	122.500	1.00 43.06	В	C
	ATOM	2049	CD	GLU	112	14.047	-98.808	121.624	1.00 44.55	В	С
	MOTA	2050	OE1	GLU	112	15.139	-99.130	122.139	1.00 45.11	В	0
	MOTA	2051	OE2	GLU	112	13.927	-98.497	120.419	1.00 46.10	В	0
30	ATOM	2052	<b>C</b> .	GLU	112	12.733	-95.825	122.844	1.00 38.09	В	C
	MOTA	2053	0	GLU	112	13.030	-95.594	121.666	1.00 37.26	В	0
	MOTA	2054	N	LEU	113	11.638	-95.349	123.430	1.00 35.31	В	N
	ATOM	2055	CA	LEU	113	10.691	-94.500	122.729	1.00 33.09	В	С
	MOTA	2056	CB	LEU	113	9.499	-94.178	123.637	1.00 32.27	В	С
35	MOTA	2057	CG	LEU	113	8.480	-93.165	123.101	1.00 32.57	В	C
	MOTA	2058	CD1	LEU	113	7.983	-93.617	121.731	1.00 31.91	В	С
	ATOM	2059	CD2	LEU	113	7.325	-93.014	124.080	1.00 31.00	В	С
	ATOM	2060	С	LEU	113	11.380	-93.213	122.294	1.00 31.56	В	С
	MOTA	2061	0	LEU	113	11.268	-92.799	121.138	1.00 30.78	В	0
40	MOTA	2062	N	ILE	114	12.089	-92.586	123.227	1.00 30.06	В	N
	ATOM	2063	CA	ILE	114	12.808	-91.351	122.949	1.00 29.43	В	С
	MOTA	2064	CB	ILE	114	13.518	-90.822	124.221	1.00 28.14	В	С
	ATOM	2065	CG2	ILE	114	14.463	~89.686	123.870	1.00 27.33	В	С
	MOTA	2066		ILE	114			125.228	1.00 27.74	В	С
45	ATOM	2067	CD1		114			126.541	1.00 26.41	В	C
	ATOM	2068	C	ILE	114			121.836	1.00 29.65	В	C
	MOTA	2069	0	ILE	114			120.872	1.00 28.66	В	0
	ATOM	2070		ARG	115			121.961	1.00 30.65	В	N
	ATOM	2071	CA	ARG	115			120.947	1.00 32.09	В	С
50	ATOM	2072	СВ	ARG	115			121.319	1.00 34.79	В	C
•	ATOM	2073	CG	ARG	115			120.729	1.00 39.78	В	C
	ATOM	2074		ARG	115			121.077	1.00 43.64	В	C
	ATOM	2075		ARG	115			120.292	1.00 47.79	В	N
	ATOM	2076		ARG	115			120.612		В	C
55	ATOM	2077		ARG	115			121.713		В	N
30	MOTA	2078		ARG	115			119.832		В	
	ATOM	2079		ARG	115			119.562		В	
	MOTA	2080		ARG				5 118.571		В	
	AT OH	2000	9			,				ט	0

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	MOTA	2081	N	THR	116	13.952	-93.734	119.496	1.00 29.49	В	N
	ATOM	2082	CA	THR	116	13.263	-93.974	118.231	1.00 28.65	В	С
	MOTA	2083	CB	THR	116	12.058	-94.914	118.428	1.00 29.45	В	С
	ATOM	2084	OG1	THR	116	12.514	-96.168	118.945	1.00 31.34	В	0
5	MOTA	2085	CG2	THR	116	11.332	-95.147	117.112	1.00 29.40	В	С
	MOTA	2086	С	THR	116	12.757	-92.658	117.640	1.00 27.59	В	С
	ATOM	2087	0	THR	116	12.995	-92.359	116.469	1.00 27.35	В	0
	ATOM	2088	N	LEU	117	12.049	-91.882	118.455	1.00 25.67	В	N
	ATOM	2089	CA	LEU	117	11.517	-90.594	118.019	1.00 24.51	В	С
10	ATOM	2090	СВ	LEU	117	10.691	-89.949	119.143	1.00 22.66	В	C
	MOTA	2091	CG	LEU	117		-90.544		1.00 23.35	В	Ċ
	ATOM	2092	CD1	LEU	117			120.731	1.00 22.38	В	Ċ
	ATOM	2093	CD2		117			118.270	1.00 22.82	В	Č
	ATOM	2094	С	LEU	117			117.607	1.00 22.88	В	Ċ
15	MOTA	2095	0	LEU	117			116.580	1.00 22.11	В	ō
	ATOM	2096	N	LEU	118			118.416	1.00 23.09	В	N
	ATOM	2097	CA	LEU	118			118.152	1.00 23.66	В	C
	MOTA	2098	СВ	LEU	118			119.276	1.00 25.11	В	Č
	ATOM	2099	CG	LEU	118			119.697	1.00 27.59	В	Ċ
20	ATOM	2100	CD1	LEU	118			120.242	1.00 28.18	В	Č
	ATOM	2101		LEU	118			118.527	1.00 28.59	В	č
	ATOM	2102	C	LEU	118			116.827	1.00 23.33	В	Č
	ATOM	2103	0	LEU	118			116.054	1.00 22.58	В	ō
	ATOM	2104	N	GLY	119			116.585	1.00 22.04	В	N
25	ATOM	2105	CA	GLY	119			115.362	1.00 21.87	В	C
	ATOM	2106	С	GLY	119			114.127	1.00 21.81	В	č
	ATOM	2107	0	GLY	119			113.191	1.00 21.52	В	ŏ
	ATOM	2108	N	ALA	120			114.122	1.00 21.37	В	N
	ATOM	2109	CA	ALA	120			113.004	1.00 21.05	В	c
30	ATOM	2110	СВ	ALA	120			113.233	1.00 20.81	В	Ċ
	MOTA	2111	С	ALA	120			112.833	1.00 21.40	В	C
	ATOM	2112	0	ALA	120			111.716	1.00 20.48	В	ō
	ATOM	2113	N	HIS	121			113.945	1.00 22.14	В	N
	ATOM.	2114	CA	HIS	121			113.914	1.00 22.47	В	C
35	MOTA	2115	СВ	HIS	121			115.325	1.00 22.23	В	C
	MOTA	2116	CG	HIS	121			115.433	1.00 24.18	В	C
	ATOM	2117	CD2	HIS	121			115.284	1.00 23.87	В	С
	ATOM	2118	ND1	HIS	121			115.657	1.00 24.73	В	N
	ATOM	2119	CE1	HIS	121			115.638	1.00 25.53	В	C
40	ATOM	2120		HIS	121			115.413	1.00 25.90	В	N
	MOTA	2121	С	HIS	121			113.339	1.00 22.36	В	С
	MOTA	2122	0	HIS	121			112.448	1.00 22.60	В	ō
	ATOM	2123	N	THR	122			113.849	1.00 21.59	В	N
	MOTA	2124	CA	THR	122	16.900	-86.119	113.387	1.00 21.97	В	C
45	MOTA	2125	СВ	THR	122			114.199	1.00 22.09	В	C
	MOTA	2126		THR	122			115.561	1.00 24.17	В	o
	MOTA	2127		THR	122			113.655	1.00 22.21	В	C
	MOTA	2128	С	THR	122			111.907	1.00 21.55	В	Č
	MOTA	2129	0	THR	122	17.664	-85.549	111.188	1.00 20.90	В	Ō
50	MOTA	2130	N	ARG	123			111.455	1.00 21.32	В	N
	ATOM	2131	CA	ARG	123	16.990	-87.969	110.063	1.00 22.23	В	C
	ATOM	2132	CB	ARG	123			109.852	1.00 21.90	В	C
	ATOM	2133	CG	ARG	123			110.237	1.00 22.07	В	C
	ATOM	2134	CD	ARG	123			109.753	1.00 20.33	В	C
55	ATOM	2135	NE	ARG	123			110.380	1.00 21.86	В	N
	ATOM	2136	CZ	ARG	123			111.601	1.00 21.57	В	C
	ATOM	2137		ARG	123			112.352	1.00 21.63	В	N
	ATOM	2138		ARG	123			112.086	1.00 20.21	В	N
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	MOTA	2139	С	ARG	123		-87.193		1.00 22.64	В	С
	MOTA	2140	0	ARG	123		-86.867		1.00 23.15	В	0
	MOTA	2141	N	HIS	124		-86.877		1.00 22.90	В	N
_	MOTA	2142	CA	HIS	124		-86.204		1.00 23.53	В	C
5	MOTA	2143	CB	HIS	124		-87.089		1.00 22.74	В	C
	ATOM	2144	CG	HIS	124		-88.501		1.00 23.34	В	C
	MOTA	2145	CD2		124			106.650	1.00 22.54	В	С
	MOTA	2146	ND1		124			108.624	1.00 22.99	В	N
10	ATOM	2147	CE1		124			108.055	1.00 21.00	В	C
10	ATOM	2148	NE2		124			106.858	1.00 21.93	В	N
	MOTA	2149 2150	C O	HIS	124			108.653 107.700	1.00 23.63 1.00 24.39	В	C
	MOTA	2150		HIS MET	124 125			107.700	1.00 24.39	В	0
	MOTA MOTA	2151	N CA		125			110.197		В	N
15	ATOM	2152	CB	MET MET	125			111.038	1.00 22.76 1.00 22.21	В	C C
13	ATOM	2154	CG	MET	125			110.415		В	С
	ATOM	2154	SD	MET	125			111.269	1.00 21.83 1.00 22.85	B B	S
	ATOM	2156	CE	MET	125			112.900	1.00 22.85	В	C
	ATOM	2157	CE	MET	125			110.880	1.00 22.78	В	C
20	ATOM	2158	Ö	MET	125			110.530	1.00 22.78	В	0
20	ATOM	2159	N	GLY	126			111.852	1.00 22.22	В	N
	ATOM	2160	CA	GLY	126			112.614	1.00 22.35	В	C
	ATOM	2161	C	GLY	126			111.919	1.00 22.45	В	C
	ATOM	2162	Ö	GLY	126			112.438	1.00 23.13	В	õ
25	ATOM	2163	N	THR	127			110.757	1.00 23.04	В	N
	ATOM	2164	CA	THR	127			110.737	1.00 23.87	В	C
	ATOM	2165	СВ	THR	127			109.920	1.00 25.68	В	C
	MOTA	2166	OG1		127			109.569	1.00 27.25	В	ŏ
	ATOM	2167	CG2		127			111.242	1.00 26.03	В	c
30	ATOM	2168	C	THR	127			108.624	1.00 22.94	В	c
	ATOM	2169	Ö	THR	127			107.728	1.00 22.72	В	ŏ
	ATOM	2170	N	MET	128			108.429	1.00 22.41	В	N
	ATOM	2171	CA	MET	128			107.114	1.00 22.16	В	C
	MOTA	2172	СВ	MET	128			107.101	1.00 21.38	В	C
35	MOTA	2173	CG	MET	128			107.943	1.00 20.15	В	C
	MOTA	2174	SD	MET	128	11.239	-80.338	107.749	1.00 20.67	В	S
	ATOM	2175	CE	MET	128	10.284	-79.362	108.917	1.00 22.56	В	С
	MOTA	2176	С	MET	128	15.305	-78.305	106.669	1.00 22.32	В	С
	MOTA	2177	0	MET	128	15.261	-78.019	105.476	1.00 22.98	В	0
40	ATOM	2178	N	PHE	129	15.363	-77.384	107.627	1.00 22.69	В	N
	MOTA	2179	CA	PHE	129			107.311	1.00 24.34	В	С
	MOTA	2180	CB	PHE	129			108.605		В	C
	ATOM	2181	CG	PHE	129			109.396	1.00 28.48	В	C
	MOTA	2182		PHE	129			109.052	1.00 30.09	В	С
45	ATOM	2183		PHE	129			110.486	1.00 29.53	В	С
	ATOM	2184		PHE	129			109.784	1.00 31.38	В	C
	MOTA	2185	CE2		129			111.228	1.00 31.21	В	C
	ATOM	2186	CZ	PHE	129			110.875	1.00 32.22	В	С
	ATOM	2187	С	PHE	129			106.485	1.00 24.09	В	С
50	MOTA	2188	0	PHE	129			105.821	1.00 23.81		0
	ATOM	2189	N	GLU	130			106.522	1.00 24.21		N
	ATOM	2190	CA	GLU	130			105.762	1.00 25.33	В	C
	ATOM	2191	CB	GLU	130			106.222	1.00 27.28	В	C
55	MOTA	2192	CG	GLU	130			3 107.685	1.00 31.35		C
55	MOTA	2193	CD	GLU	130			7 108.045	1.00 34.28		C
	ATOM	2194		GLU	130			107.389	1.00 35.08		0
	ATOM	2195		GLU	130			3 108.992	1.00 36.90		0
	MOTA	2196	С	GLU	130	18.647	-/6.19	7 104.258	1.00 23.24	В	C

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	MOTA	2197	0	GLU	130	19.439	-75.686	103.470	1.00 23.56	В	0
	ATOM	2198	N	GLN	131	17.601	-76.918	103.865	1.00 21.55	В	N
	MOTA	2199	CA	GLN	131	17.302	-77.125	102.453	1.00 20.76	В	С
	ATOM	2200	СВ	GLN	131	16.539	-78.442	102.240	1.00 21.85	В	С
5	MOTA	2201	CG	GLN	131	17.320	-79.703	102.536	1.00 23.34	В	С
	ATOM	2202	CD	GLN	131	18.691	-79.696	101.882	1.00 26.07	В	С
	ATOM	2203	OE1	GLN	131		-79.538		1.00 26.66	В	0
	ATOM	2204		GLN	131		-79.862		1.00 26.55	В	N
	ATOM	2205	С	GLN	131		-75.987		1.00 20.02	В	C
10	ATOM	2206	0	GLN	131		-75.875		1.00 20.04	В	Ō
	ATOM	2207	N	PHE	132		-75.149		1.00 19.42	В	N
	ATOM	2208	CA	PHE	132		-74.048		1.00 19.05	В	C
	ATOM	2209	СВ	PHE	132		-73.157		1.00 17.06	В	Ċ
	ATOM	2210	CG	PHE	132		-73.779		1.00 16.44	В	Č
15	ATOM	2211		PHE	132		-75.055		1.00 15.33	В	c
	ATOM	2212	CD2		132		-73.073		1.00 15.55	В	c
	ATOM	2213	CE1		132		-75.620		1.00 15.93	В	C
	MOTA	2214	CE2	PHE	132		-73.626		1.00 15.23	В	C
	ATOM	2215	CZ	PHE	132		-74.895		1.00 15.25	В	C
20	ATOM	2216	C	PHE	132		-73.182		1.00 13.13	В	C
20	ATOM	2217	Ö	PHE	132		-72.736		1.00 19.75	В	0
	ATOM	2218	N	VAL	133		-72.736				
		2219	CA	VAL	133		-72.346		1.00 20.25	В	N
	MOTA	2220	CB	VAL	133				1.00 21.53	В	C
25	MOTA			VAL			-71.915		1.00 22.78	В	С
25	ATOM	2221		VAL	133		-73.236		1.00 21.70	В	C
	ATOM	2222			133		-70.870	99.850	1.00 23.73	В	C
	ATOM	2223	C	VAL	133		-72.750	98.877	1.00 22.37	В	С
	ATOM	2224	0	VAL	133		-72.058	97.895	1.00 21.91	В	0
20	MOTA	2225	N	GLN	134		-74.036	98.780	1.00 22.29	В	N
30	MOTA	2226	CA	GLN	134		-74.725	97.496	1.00 23.37	В	С
	MOTA	2227	CB	GLN	134		-76.229		1.00 22.92	В	С
	MOTA	2228	CG	GLN	134		-76.621	98.407	1.00 23.40	В	C
	MOTA	2229	CD	GLN	134		-76.250		1.00 23.05	В	C
0.5	ATOM	2230	OE1		134		-76.701	98.020	1.00 25.27	В	0
35	MOTA	2231		GLN	134		-75.430		1.00 22.55	В	N
	ATOM	2232	С	GLN	134		-74.548		1.00 24.55	В	С
	ATOM	2233	0	GLN	134		~75.081		1.00 24.39	В	0
	ATOM	2234	N	PHE	135		-73.804		1.00 24.64	В	N
	MOTA	2235	CA	PHE	135	13.710	-73.595	96.750	1.00 25.44	В	С
40	ATOM	2236	CB	PHE.	135		-74.065		1.00 24.79	В	C
	MOTA	2237	CG	PHE	135	12.706	~75.534	98.039	1.00 23.89	В	C
	MOTA	2238	CD1	PHE	135	12.206	-76.478	97.146	1.00 24.40	В	С
	MOTA	2239	CD2	PHE	135	13.335	-75.977	99.198	1.00 24.10	В	С
	MOTA	2240	CE1	PHE	135	12.331	-77.848	97.397	1.00 24.28	В	С
45	MOTA	2241	CE2	PHE	135	13.470	-77.349	99.466	1.00 24.09	В	С
	MOTA	2242	CZ	PHE	135	12.966	-78.286	98.560	1.00 24.17	В	С
	MOTA	2243	С	PHE	135	13.500	-72.130	96.358	1.00 27.05	В	С
	MOTA	2244	0	PHE	135	12.508	-71.501		1.00 27.09	В	0
	ATOM	2245	N	ARG	136	14.444	-71.622		1.00 27.76	В	N
50	MOTA	2246	CA	ARG	136		-70.241		1.00 28.77	В	C
	ATOM	2247	CB	ARG	136	14.046	-70.118		1.00 30.48	В	C
	ATOM	2248	CG	ARG	136		-70.801		1.00 34.79	В	C
	MOTA	2249	CD	ARG			-72.190		1.00 36.21	В	C
	ATOM	2250	NE	ARG			-73.127		1.00 39.78		N
55	MOTA	2251	CZ	ARG			-74.448		1.00 40.31	В	C
	MOTA	2252		ARG			-75.011		1.00 41.22		N
	MOTA	2253	NH2				-75.207		1.00 40.63		
	ATOM	2254	C	ARG			-69.204				
	ATOM	2234	C	MRG	120	13./05	-09.204	33.331	1.00 28.40	B	С

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	ATOM	2255	0	ARG	136	12.679		95.618	1.00 27.33	В	0
	MOTA	2256	N	PRO	137	14.379		97.085	1.00 28.01	В	N
	ATOM	2257	CD	PRO	137	15.580		97.667	1.00 27.43	В	С
_	MOTA	2258	CA	PRO	137		-67.860	97.999	1.00 27.65	В	C
5	MOTA	2259	CB	PRO	137		-68.120	99.286	1.00 27.04	В	С
	MOTA	2260	CG	PRO	137		-68.518	98.778	1.00 27.95	В	С
	MOTA	2261	С	PRO	137		-66.467	97.451	1.00 27.46	В	С
	MOTA	2262	0	PRO	137		-66.190	96.981	1.00 27.51	В	0
4.0	MOTA	2263	N	PRO	138		-65.578	97.479	1.00 26.95	В	N
10	MOTA	2264	CD	PRO	138		-65.734	97.852	1.00 26.96	В	C
	MOTA	2265	CA	PRO	138		-64.234	96.968	1.00 26.24	В	С
	MOTA	2266	CB	PRO	138		-63.496	97.228	1.00 26.54	В	С
	MOTA	2267	CG	PRO	138		-64.341	98.273	1.00 27.40	В	С
45	MOTA	2268	С	PRO	138		-63.608	97.679	1.00 25.08	В	C
15	MOTA	2269	0	PRO	138		-63.992	98.799	1.00 23.40	В	0
	MOTA	2270	N	ALA	139		-62.642	97.014	1.00 24.45	В	N
	MOTA	2271	CA	ALA	139		-61.959	97.529	1.00 24.74	В	С
	MOTA	2272	CB	ALA	139		-60.925	96.503	1.00 25.14	В	C
00	MOTA	2273	С	ALA	139		-61.300	98.905	1.00 24.49	В	С
20	MOTA	2274	0	ALA	139		-61.293	99.651	1.00 24.23	В	0
	ATOM	2275	N	HIS	140		-60.749	99.252	1.00 24.55	В	N
	MOTA	2276	CA	HIS	140		-60.066		1.00 24.29	В	С
	MOTA	2277	СВ	HIS	140		-59.245		1.00 23.87	В	C
05	MOTA	2278	CG	HIS	140		-60.056		1.00 23.38	В	С
25	MOTA	2279		HIS	140		-60.232		1.00 23.33	В	C
	ATOM	2280		HIS	140		-60.783	99.833	1.00 23.88	В	N
	MOTA	2281		HIS	140		-61.371		1.00 24.02	В	С
	ATOM	2282		HIS	140		-61.054		1.00 24.40	В	N
20	ATOM	2283	C	HIS	140		-60.951		1.00 24.34	В	C
30	ATOM	2284	0	HIS	140		-60.439		1.00 24.05	В	0
	ATOM	2285	N	LEU	141		-62.265		1.00 24.45	В	N
	ATOM	2286	CA	LEU	141		-63.216		1.00 25.64	В	C
	ATOM	2287	CB	LEU	141		-64.544		1.00 24.52	В	C
35	ATOM	2288	CG	LEU	141		-64.562		1.00 25.14	В	C
35	ATOM	2289		LEU LEU	141		-65.992		1.00 23.52	В	C
	ATOM	2290			141		-64.006		1.00 23.73	В	C
	ATOM	2291	C	LEU	141		-63.512		1.00 26.62	В	C
	ATOM ATOM	2292 2293	O	LEU	141		-64.005	104.138	1.00 26.38	В	0
40	ATOM	2294	N CA	PHE	142 142		-63.495		1.00 28.49	В	N
70	ATOM	2295	CB	PHE	142		-63.519		1.00 31.82	В	C
	ATOM	2296			142			100.985	1.00 29.28 1.00 27.30	В	
	ATOM	2297		PHE	142		-65.873		1.00 27.30		
	ATOM	2298		PHE	142		-64.890	99.331	1.00 25.11	B B	C
45	ATOM	2299		PHE	142		-67.056	99.706	1.00 25.73	В	C
70	ATOM	2300		PHE	142		-66.074	98.639	1.00 25.73	В	C
	ATOM	2301	CZ	PHE	142		-67.159	98.826	1.00 25.25	В	C
	ATOM	2302	C	PHE	142			103.275	1.00 25.25	В	C
	ATOM	2303	Ö	PHE	142			103.273	1.00 35.28	В	o
50	ATOM	2304	N	ILE	143			103.201	1.00 33.37	В	N
00	ATOM	2305	CA	ILE	143			105.028	1.00 33.02	В	C
	ATOM	2306	CB	ILE	143			103.028	1.00 42.74	В	C
	ATOM	2307		ILE	143			104.472	1.00 42.50	В	
	ATOM	2308		ILE	143			103.329	1.00 42.81	В	C
55	MOTA	2309	CD1		143			104.032	1.00 42.81	В	C
	MOTA	2310	C	ILE	143			105.388	1.00 45.32	В	C
	ATOM	2311	Ö	ILE	143			106.333	1.00 45.22	В	0
	ATOM	2312	N	HIS	144			104.636	1.00 48.12	В	
	AIOM	2312	**		743	21.402	50.105	104.030	1.00 40.10	8	N

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	ATOM	2313	CA	HIS	144	21.138	-58.766	104.905	1.00 50.43	В	C
	MOTA	2314	CB	HIS	144	22.347	-58.077	105.549	1.00 51.71	В	C
	ATOM	2315	CG	HIS	144	22.639	-58.553	106.941	1.00 53.18	В	С
	MOTA	2316	CD2	HIS	144	21.840	-59.129	107.873	1.00 53.68	В	С
5	MOTA	2317	ND1	HIS	144	23.885	-58.447	107.520	1.00 53.87	В	N
	MOTA	2318	CE1	HIS	144	23.843	-58.939	108.747	1.00 53.89	В	C
	MOTA	2319	NE2	HIS	144	22.613	-59.360	108.985	1.00 53.40	В	N
	MOTA	2320	С	HIS	144	20.688	-58.010	103.655	1.00 50.90	В	С
	MOTA	2321	0	HIS	144	21.479	-57.322	103.006	1.00 51.65	В	0
10	MOTA	2322	N	HIS	145	19.403	-58.149	103.339	1.00 51.36	В	N
	MOTA	2323	CA	HIS	145	18.778	-57.503	102.183	1.00 51.48	В	С
	MOTA	2324	CB	HIS	145	18.251	-58.577	101.223	1.00 52.46	В	C
	ATOM	2325	CG	HIS	145	18.066	-58.105	99.812	1.00 53.73	В	С
	MOTA	2326	CD2	HIS	145	16.958	-57.714	99.140	1.00 54.04	В	С
15	ATOM	2327	ND1	HIS	145	19.109	-58.015	98.915	1.00 54.94	В	N
	ATOM	2328	CE1	HIS	145	18.651	-57.592	97.750	1.00 54.82	В	C
	MOTA	2329	NE2	HIS	145	17.348	-57.401	97.859	1.00 54.54	В	N
	ATOM	2330	С	HIS	145	17.610	-56.629	102.673	1.00 50.58	В	С
	MOTA	2331	0	HIS	145	16.980	-56.929	103.691	1.00 50.62	В	0
20	MOTA	2332	N	GLN	146	17.325	-55.552	101.950	1.00 49.68	В	N
	MOTA	2333	CA	GLN	146	16.227	-54.650	102.303	1.00 48.58	В	С
	MOTA	2334	CB	GLN	146	16.264	-53.426	101.373	1.00 50.29	В	C
	ATOM	2335	CG	GLN	146	15.242	-52.329	101.656	1.00 52.37	В	С
	ATOM	2336	CD	GLN	146	14.090	-52.316	100.661	1.00 53.97	В	C
25	ATOM	2337	OE1	GLN	146	14.299	-52.395	99.446	1.00 55.36	В	0
	MOTA	2338	NE2	GLN	146	12.867	-52.216	101.172	1.00 53.39	В	N
	ATOM	2339	С	GLN	146	14.902	-55.412	102.152	1.00 46.43	В	С
	MOTA	2340	0	GLN	146	14.772	-56.250	101.259	1.00 47.01	В	0
	ATOM	2341	N	PRO	147	13.906	-55.140	103.024	1.00 43.87	В	N
30	MOTA	2342	CD	PRO	147	13.896	-54.126	104.094	1.00 42.94	В	C
	MOTA	2343	CA	PRO	147	12.604	-55.823	102.952	1.00 40.85	В	С
	MOTA	2344	CB	PRO	147	11.805	-55.171	104.081	1.00 41.31	В	С
	ATOM	2345	CG	PRO	147	12.427	-53.812	104.211	1.00 42.30	В	С
	MOTA	2346	С	PRO	147	11.911	-55.697	101.594	1.00 37.44	В	C
35	ATOM	2347	0	PRO	147	12.368	-54.956	100.733	1.00 36.62	В	0
	ATOM	2348	N	LEU	148	10.812	-56.428	101.408	1.00 34.45	В	N
	ATOM	2349	CA	LEU	148	10.076	-56.402	100.143	1.00 31.79	В	C
	ATOM	2350	CB	LEU	148	8.912	-57.397	100.162	1.00 30.82	В	C
	MOTA	2351	CG	LEU	148		-58.528		1.00 31.47	В	С
40	ATOM	2352		. LEU	148	7.437	-58.918	98.878	1.00 30.58	В	С
	MOTA	2353	CD2	LEU	148	9.531	-58.102	97.817	1.00 30.15	В	С
	MOTA	2354	С	LEU	148	9.521	-55.018	99.849	1.00 29.43	В	С
	MOTA	2355	0	LEU	148	8.779	-54.456	100.653	1.00 28.77		0
	ATOM	2356	N	PRO	149		-54.448		1.00 27.64	В	N
45	MOTA	2357	CD	PRO	149	10.846	-54.900	97.686	1.00 27.10	В	C
	MOTA	2358	CA	PRO	149	9.358	-53.114	98.361	1.00 26.65	В	
	ATOM	2359	СВ	PRO	149	9.986	-52.829	96.998	1.00 26.71	В	
	ATOM	2360	CG	PRO	149	11.296	-53.591	. 97.075	1.00 26.43	В	
	ATOM	2361	С	PRO	149	7.833	-53.103	98.311	1.00 25.49	В	C
50	ATOM	2362	0	PRO	149	7.201	54.132	98.090	1.00 25.29	В	0
	MOTA	2363	N	THR	150	7.248	-51.932	98.516	1.00 24.33	В	N
	ATOM	2364	CA	THR	150	5.799	-51.782	98.494	1.00 23.46	В	C
	ATOM	2365	СВ	THR	150	5.417	-50.287	98.643	1.00 23.36	В	
	MOTA	2366	OG1	L THR	150	5.760	-49.841	99.964	1.00 21.71	В	
55	MOTA	2367	CG2	2 THR		3.934	-50.07	7 98.395	1.00 22.73	В	
	ATOM	2368	С	THR	150	5.120	-52.35	7 97.244	1.00 23.83	В	
	MOTA	2369		THR			-53.050				
	MOTA	2370	N	LEU	151	5.671	-52.07	96.071	1.00 23.39	В	N

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	MOTA	2371	CA	LEU	151		-52.549	94.818	1.00 24.18	В	С
	MOTA	2372	CB	LEU	151	5.174	-51.435	93.764	1.00 23.98	В	С
	MOTA	2373	CG	LEU	151		-50.542	93.516	1.00 26.19	В	С
_	MOTA	2374	CD1		151		-50.352	94.783	1.00 25.29	В	С
5	MOTA	2375	CD2	LEU	151		-49.199	92.956	1.00 25.33	В	С
	MOTA	2376	С	LEU	151	5.692	-53.843	94.256	1.00 24.80	В	С
	MOTA	2377	0	LEU	151		-54.307	93.188	1.00 24.09	В	0
	MOTA	2378	N	ALA	152		-54.433	94.965	1.00 25.14	В	N
	MOTA	2379	CA	ALA	152		-55.666	94.485	1.00 25.42	В	С
10	MOTA	2380	СВ	ALA	152		-56.032	95.357	1.00 25.27	В	С
	MOTA	2381	С	ALA	152		-56.800	94.483	1.00 25.45	В	С
	MOTA	2382	0	ALA	152		-56.990	95.447	1.00 26.33	В	0
	MOTA	2383	N	PRO	153		-57.558	93.385	1.00 25.59	В	N
	MOTA	2384	CD	PRO	153		-57.411	92.078	1.00 26.03	В	С
15	MOTA	2385	CA	PRO	153		-58.658	93.373	1.00 25.15	В	С
	MOTA	2386	СВ	PRO	153		-59.222	91.953	1.00 25.93	В	C
	ATOM	2387	CG	PRO	153		-58.795	91.514	1.00 27.37	В	С
	MOTA	2388	С	PRO	153		-59.683	94.457	1.00 24.29	В	С
	MOTA	2389	0	PRO	153		-59.912	94.776	1.00 23.77	В	0
20	MOTA	2390	N	VAL	154		-60.283	95.031	1.00 23.14	В	N
	MOTA	2391	CA	VAL	154	4.689	-61.269	96.090	1.00 23.20	В	С
	MOTA	2392	CB	VAL	154		-61.118	97.205	1.00 23.14	В	С
	MOTA	2393		VAL	154	3.638	-59.705	97.761	1.00 24.25	В	С
~=	MOTA	2394		VAL	154		-61.440	96.656	1.00 22.99	В	С
25	MOTA	2395	С	VAL	154		-62.710	95.589	1.00 22.02	В	С
	MOTA	2396	0	VAL	154		-63.631	96.364	1.00 21.64	В	0
	MOTA	2397	N	LEU	155		-62.901	94.298	1.00 21.95	В	N
	ATOM	2398	CA	LEU	155	4.374	-64.244	93.711	1.00 21.77	В	C
	MOTA	2399	CB	LEU	155	4.170	-64.164	92.191	1.00 21.08	В	С
30	MOTA	2400	CG	LEU	155	4.166	-65.490	91.410	1.00 22.05	В	С
	MOTA	2401		LEU	155		-66.385	91.892	1.00 20.48	В	С
	MOTA	2402		LEU	155		-65.204	89.905	1.00 22.31	В	C
	MOTA	2403	С	LEU	155		-65.083	94.030	1.00 20.95	В	С
0.5	MOTA	2404	0	LEU	155		-66.258	94.373	1.00 21.26	В	0
35	MOTA	2405	N	PRO	156		-64.505	93.908	1.00 20.25	В	N
	MOTA	2406	CD	PRO	156		-63.205	93.313	1.00 20.55	В	С
	MOTA	2407	CA	PRO	156		-65.294	94.218	1.00 20.02	В	С
	MOTA	2408	CB	PRO	156		-64.316	93.942	1.00 19.60	В	С
40	MOTA	2409	CG	PRO	156		-63.462	92.821	1.00 19.34	В	С
40	MOTA	2410	С	PRO	156		-65.793	95.675	1.00 20.44	В	С
	MOTA	2411	0	PRO	156		-66.943	95.948	1.00 18.86	В	0
	MOTA	2412	N	LEU	157		-64.922	96.603	1.00 18.95	В	N
	MOTA	2413	CA	LEU	157		-65.298	98.012	1.00 18.72	В	С
45	MOTA	2414	CB	LEU	157		-64.086	98.886	1.00 18.06	В	C
45	MOTA	2415	CG	LEU	157			100.390	1.00 18.20	В	С
	MOTA	2416		LEU	157		-64.931	100.910	1.00 18.30	В	С
	MOTA	2417		LEU	157			101.133	1.00 18.86	В	C
	MOTA	2418	С	LEU	157		-66.376	98.210	1.00 18.16	В	С
	MOTA	2419	0	LEU	157		-67.369	98.892	1.00 17.47	В	0
50	MOTA	2420	N	VAL	158		-66.174	97.612	1.00 17.84	В	N
	MOTA	2421	CA	VAL	158		-67.132	97.713	1.00 18.75	В	С
	MOTA	2422	СВ	VAL	158		-66.637	96.928	1.00 19.19	В	С
	ATOM	2423		VAL	158		-67.746	96.840	1.00 18.53	В	С
	MOTA	2424		VAL	158		-65.397	97.610	1.00 19.01	В	С
55	ATOM	2425	С	VAL	158		-68.504	97.167	1.00 19.26	В	С
	ATOM	2426	0	VAL	158		-69.539	97.773	1.00 17.72	В	0
	ATOM	2427	N	THR	159		-68.504	96.009	1.00 19.18	В	N
	ATOM	2428	CA	THR	159	5.749	-69.742	95.382	1.00 20.12	В	С

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	MOTA	2429	СВ	THR	159	6.359	-69.464	94.005	1.00 20.73	В	С
	MOTA	2430	OG1	THR	159	5.429	-68.696	93.233	1.00 21.98	В	0
	MOTA	2431	CG2	THR	159	6.670	-70.780	93.277	1.00 20.98	В	С
_	MOTA	2432	C	THR	159	6.786	-70.418	96.270	1.00 19.12	В	С
5	MOTA	2433	0	THR	159	6.757	-71.631	96.447	1.00 19.81	В	0
	MOTA	2434	N	HIS	160	7.695	-69.624	96.827	1.00 18.60	В	N
	MOTA	2435	CA	HIS	160	8.725	-70.143	97.722	1.00 19.62	В	С
	MOTA	2436	CB	HIS	160	9.628	-69.004	98.201	1.00 19.12	В	С
	MOTA	2437	CG	HIS	160	10.583	-69.405	99.283	1.00 20.97	В	С
10	MOTA	2438	CD2	HIS	160	10.739	-68.946	100.548	1.00 19.99	В	С
	MOTA	2439	ND1	HIS	160	11.547	-70.374	99.106	1.00 21.00	В	N
	MOTA	2440	CE1	HIS	160	12.258	-70.492	100.214	1.00 20.06	В	С
	ATOM	2441	NE2	HIS	160	11.787	-69.637	101.103	1.00 18.80	В	N
	MOTA	2442	С	HIS	160	8.069	-70.835	98.922	1.00 18.85	В	С
15	MOTA	2443	0	HIS	160	8.465	-71.934	99.309	1.00 19.48	В	0
	MOTA	2444	N	PHE	161	7.065	-70.192	99.510	1.00 18.30	В	N
	MOTA	2445	CA	PHE	161	6.362	-70.783	100.644	1.00 17.81	В	С
	MOTA	2446	CB	PHE	161	5.333	-69.799	101.218	1.00 17.09	В	С
	MOTA	2447	CG	PHE	161	5.924	-68.773	102.167	1.00 18.77	В	С
20	MOTA	2448	CD1	PHE	161	7.253	-68.861	102.576	1.00 18.08	В	C
	MOTA	2449	CD2	PHE	161	5.137	-67.743	102.675	1.00 18.45	В	С
	MOTA	2450	CE1	PHE	161	7.788	-67.943	103.478	1.00 19.86	В	С
	ATOM	2451	CE2	PHE	161	5.658	-66.818	103.577	1.00 19.78	В	С
	MOTA	2452	CZ	PHE	161	6.992	-66.918	103.981	1.00 20.06	В	С
25	MOTA	2453	С	PHE	161	5.675	-72.081	100.227	1.00 17.47	В	С
	MOTA	2454	0	PHE	161	5.721	-73.075	100.953	1.00 16.02	В	0
	ATOM	2455	N	ALA	162	5.039	-72.082	99.058	1.00 16.64	В	N
	ATOM	2456	CA	ALA	162	4.361	-73.286	98.588	1.00 17.85	В	С
	MOTA	2457	CB	ALA	162	3.712	-73.034	97.224	1.00 17.30	В	С
30	ATOM	2458	С	ALA	162	5.352	-74.447	98.487	1.00 18.28	В	С
	MOTA	2459	0	ALA	162	5.055	-75.563	98.899	1.00 17.59	В	0
	ATOM	2460	N	ASP	163	6.530	-74.166	97.935	1.00 18.81	В	N
	MOTA	2461	CA	ASP	163	7.569	-75.170	97.763	1.00 20.29	В	С
	MOTA	2462	CB	ASP	163	8.650	-74.624	96.834	1.00 21.76	В	С
35	ATOM	2463	CG	ASP	163	8.147	-74.448	95.414	1.00 22.93	В	С
	ATOM	2464	OD1	ASP	163	8.816	-73.763	94.621	1.00 25.09	В	0
	MOTA	2465	OD2	ASP	163	7.080	-75.005	95.093	1.00 23.14	В	0
	ATOM	2466	С	ASP	163	8.183	-75.671	99.072	1.00 20.12	В	C
	ATOM	2467	0	ASP	163	8.307	-76.873	99.258	1.00 18.59	В	0
40	ATOM	2468	N	ILE	164	8.564	-74.780	99.986	1.00 19.96	В	N
	MOTA	2469	CA	ILE	164	9.117	-75.286	101.240	1.00 20.35	В	С
	ATOM	2470	CB	ILE	164	9.877	-74.193	102.071	1.00 20.04	В	С
	MOTA	2471	CG2	ILE	164	11.003	-73.596	101.224	1.00 19.12	В	С
	MOTA	2472	CG1	ILE	164	8.934	-73.093	102.555	1.00 20.01	В	C
45	MOTA	2473	CD1	ILE	164	9.619	-72.108	103.497	1.00 18.11	В	C
	ATOM	2474	C	ILE	164	8.014	-75.950	102.076	1.00 20.60	В	С
	ATOM	2475	0	ILE	164	8.287	-76.890	102.819	1.00 20.54	В	0
	MOTA	2476	N	ASN	165	6.766	-75.496	101.939	1.00 19.91	В	N
	ATOM	2477	CA	ASN	165	5.664	-76.123	102.673	1.00 19.83	В	С
50	ATOM	2478	CB	ASN	165	4.341	-75.370	102.464	1.00 18.46	В	С
	ATOM	2479	CG	ASN	165			103.244	1.00 19.03	В	С
	MOTA	2480		ASN	165	5.070	-73.812	104.145	1.00 17.31	В	0
	ATOM	2481	ND2	ASN	165			102.900	1.00 17.09	В	N
	ATOM	2482	С	ASN	165			102.174	1.00 20.04	В	С
55	MOTA	2483	0	ASN	165	5.334	-78.500	102.960	1.00 19.45	В	0
	MOTA	2484	N	THR	166	5.525	-77.741	100.859	1.00 20.25	В	N
	MOTA	2485	CA	THR	166	5.379	-79.071	100.279		В	С
	ATOM	2486	CB	THR	166		-78.981			В	С

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	MOTA	2487	OG1	THR	166	4.211	-78.224	98.341	1.00 22.00	В	0
	MOTA	2488	CG2	THR	166		-80.368	98.104	1.00 20.43	В	С
	MOTA	2489	С	THR	166		-79.947		1.00 20.47	В	C
_	MOTA	2490	Ο.	THR	166	6.338	-81.079	101.198	1.00 19.94	В	0
5	MOTA	2491	N	PHE	167	7.745	-79.403	100.682	1.00 19.96	В	N
	MOTA	2492	CA	PHE	167		-80.099		1.00 20.53	В	С
	MOTA	2493	CB	PHE	167		-79.166		1.00 20.58	В	С
	MOTA	2494	CG	PHE	167		-79.620		1.00 21.61	В	С
	MOTA	2495	CD1		167		-80.739		1.00 21.76	В	С
10	MOTA	2496	CD2		167		-78.909		1.00 20.70	В	С
	MOTA	2497	CE1	PHE	167		-81.145		1.00 22.38	В	С
	MOTA	2498	CE2	PHE	167		-79.303		1.00 22.67	В	С
	MOTA	2499	CZ	PHE	167	13.685	-80.426	103.137	1.00 22.67	В	C
	MOTA	2500	C	PHE	167		-80.557		1.00 20.25	В	C
15	MOTA	2501	0	PHE	167		-81.730		1.00 20.74	В	0
	MOTA	2502	N	MET	168		-79.640		1.00 18.80	В	N
	MOTA	2503	CA	MET	168	8.235	-79.971	104.897	1.00 18.84	В	С
	MOTA	2504	CB	MET	168	7.858	-78.722	105.712	1.00 17.54	В	С
	MOTA	2505	CG	MET	168	9.004	-77.727	105.904	1.00 16.54	В	С
20	ATOM	2506	SD	MET	168	8.609	-76.401	107.082	1.00 16.62	В	S
	ATOM	2507	CE	MET	168	7.812	-75.178	106.019	1.00 15.56	В	C
	ATOM	2508	С	MET	168	7.215	-81.078	105.163	1.00 18.68	В	C
	MOTA	2509	0	MET	168	7.461	-81.969	105.970	1.00 17.64	В	0
	MOTA	2510	N	VAL	169	6.065	-81.024	104.503	1.00 19.45	В	N
25	ATOM	2511	CA	VAL	169	5.057	-82.064	104.702	1.00 20.76	В	С
	ATOM	2512	CB	VAL	169	3.770	-81.768	103.889	1.00 21.33	В	С
	MOTA	2513	CG1	VAL	169	2.833	-82.969	103.936	1.00 20.85	В	С
	MOTA	2514	CG2	VAL	169	3.072	-80.531	104.458	1.00 21.08	В	C
	MOTA	2515	С	VAL	169	5.642	-83.414	104.275	1.00 21.26	В	C
30	MOTA	2516	0	VAL	169	5.475	-84.420	104.965	1.00 20.54	В	0
	MOTA	2517	N	LEU	170	6.337	-83.435	103.139	1.00 21.18	В	N
	MOTA	2518	CA	LEU	170	6.959	-84.670	102.659	1.00 21.80	В	С
	MOTA	2519	CB	LEU	170	7.695	-84.421	101.338	1.00 22.86	В	С
	ATOM	2520	CG	LEU	170	6.802	-84.091	100.139	1.00 24.31	В	С
35	MOTA	2521	CD1	LEU	170	7.655	-83.710	98.941	1.00 24.24	В	C
	ATOM	2522	CD2	LEU	170	5.928	-85.300	99.818	1.00 24.97	В	C
	ATOM	2523	С	LEU	170	7.942	-85.212	103.702	1.00 21.40	В	С
	ATOM	2524	0	LEU	170			103.891	1.00 20.50	в	0
	MOTA	2525	N	GLN	171			104.371	1.00 20.25	В	N
40	ATOM	2526	CA	GLN	171			105.402	1.00 20.37	В	C
	ATOM	2527	CB	GLN	171			105.808	1.00 18.76	в	C
	ATOM	2528	CG	GLN	171	11.490	-83.112	104.719	1.00 17.74	В	
	MOTA	2529	CD	GLN	171	12.377	-84.259	104.254	1.00 18.92	В	Č
	MOTA	2530	OE1	GLN	171			105.034	1.00 20.91	В	ō
45	MOTA	2531		GLN	171			102.987	1.00 17.61	В	N
	ATOM	2532	С	GLN	171			106.626	1.00 20.12	В	C
	ATOM	2533	0	GLN	171			107.249	1.00 19.85	В	ō
	ATOM	2534	N	VAL	172			106.971	1.00 20.71	В	N
	ATOM	2535	CA	VAL	172			108.112	1.00 21.73	В	C
50	ATOM	2536	СВ	VAL	172			108.363	1.00 22.96	В	c
-	ATOM	2537		VAL	172			109.384	1.00 22.67	В	c
	ATOM	2538		VAL	172			108.874	1.00 22.87	В	C
	ATOM	2539	C	VAL	172			107.853	1.00 21.64	В	C
	ATOM	2540	ŏ	VAL	172			108.755	1.00 21.64	В	0
55	ATOM	2541	N	ILE	173			106.612	1.00 20.04	В	И
	ATOM	2542	CA	ILE	173			106.012	1.00 21.03	В	C
	ATOM	2543	CB	ILE	173			104.725	1.00 21.75	В	C
	MOTA	2544		ILE	173			104.723	1.00 21.83	В	C
					-,,	4.771	-07.303	104.2/1	1.00 22.44	D	C

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	MOTA	2545	CG1	ILE	173		-87.222		1.00 21.48	В	С
	MOTA	2546	CD1	ILE	173	3.360	-87.129	103.143	1.00 20.80	В	С
	MOTA	2547	С	ILE	173	6.725	-89.126	106.449	1.00 21.99	В	C
	MOTA	2548	0	ILE	173	6.512	-90.212	106.986	1.00 20.98	В	0
5	MOTA	2549	N	LYS	174	7.942	-88.742	106.057	1.00 22.27	В	N
	MOTA	2550	CA	LYS	174	9.117	-89.596	106.252	1.00 22.72	В	С
	MOTA	2551	CB	LYS	174	10.365	-88.925	105.671	1.00 23.87	В	С
	ATOM	2552	CG	LYS	174	10.571	-89.108	104.171	1.00 27.23	В	С
	MOTA	2553	CD	LYS	174	11.763	-88.274	103.704	1.00 29.56	В	С
10	MOTA	2554	CE	LYS	174	12.282	-88.701		1.00 32.40	В	C
	MOTA	2555	NZ	LYS	174	13.087	-89.966	102.435	1.00 34.29	В	N
	ATOM	2556	С	LYS	174	9.344	-89.888	107.743	1.00 21.87	В	С
	MOTA	2557	0	LYS	174	9.709	-91.007	108.133	1.00 20.46	В	0
	ATOM	2558	N	PHE	175	9.141	-88.862	108.563	1.00 21.53	В	N
15	ATOM	2559	CA	PHE	175	9.291	-88.966	110.008	1.00 21.67	В	С
	ATOM	2560	СВ	PHE	175	9.027	-87.599	110.643	1.00 21.39	В	С
	ATOM	2561	CG	PHE	175	8.820	-87.642	112.134	1.00 22.55	В	С
	MOTA	2562		PHE	175		-87.933		1.00 21.38	В	С
	ATOM	2563		PHE	175			112.679	1.00 22.64	В	Ċ
20	ATOM	2564		PHE	175				1.00 21.89	В	C
	ATOM	2565		PHE	175			114.061	1.00 23.00	В	Č
	ATOM	2566	CZ	PHE	175			114.910	1.00 22.35	В	Ċ
	ATOM	2567	C	PHE	175			110.581	1.00 21.68	В	c
	ATOM	2568	ō	PHE	175			111.362	1.00 21.47	В	ŏ
25	MOTA	2569	N	THR	176			110.188	1.00 22.73	В	N
	ATOM	2570	CA	THR	176			110.686	1.00 24.75	В	C
	ATOM	2571	СВ	THR	176			110.230	1.00 25.26	В	c
	ATOM	2572		THR	176			108.820	1.00 24.98	В	ŏ
	ATOM	2573	CG2		176			110.565	1.00 24.45	В	Č
30	ATOM	2574	c	THR	176			110.257	1.00 25.69	В	č
	ATOM	2575	ō	THR	176			111.065	1.00 24.44	В	ō
	MOTA	2576	N	LYS	177			109.005	1.00 26.56	В	N
	ATOM	2577	CA	LYS	177			108.489	1.00 28.90	В	C
	ATOM	2578	CB	LYS	177			106.958	1.00 29.06	В	Č
35	ATOM	2579	CG	LYS	177			106.327	1.00 31.35	В	Ċ
	ATOM	2580	CD	LYS	177	5.742		104.815	1.00 33.93	В	Ċ
	ATOM	2581	CE	LYS	177	4.452		104.191	1.00 35.45	В	Ċ
	ATOM	2582	NZ	LYS	177	3.233		104.749	1.00 36.90	В	N
	ATOM	2583	C	LYS	177	8.265		109.047	1.00 29.97	В	c
40	ATOM	2584	ō	LYS	177	8.581			1.00 30.08	В	ō
	MOTA	2585	Ŋ	ASP	178			109.924	1.00 30.95	В	N
	ATOM	2586	CA	ASP	178			110.580	1.00 31.86	В	C
	MOTA	2587	СВ	ASP	178			110.695	1.00 33.44	В	Ċ
	ATOM	2588	CG	ASP	178			109.689	1.00 35.31	В	C
45	MOTA	2589		ASP	178			108.587	1.00 36.42	В	ō
. •	MOTA	2590		ASP	178			109.994	1.00 37.36	В	Ö
	ATOM	2591	C	ASP	178			111.986	1.00 31.51	В	C
	ATOM	2592	Ō	ASP	178			112.760	1.00 31.62	В	ō
	ATOM	2593	N	LEU	179			112.299	1.00 31.35	В	N
50	ATOM	2594	CA	LEU	179			113.593	1.00 31.19	В	C
-	ATOM	2595	CB	LEU	179			114.136	1.00 31.02	В	Ċ
	ATOM	2596	CG	LEU	179			114.840	1.00 31.99	В	c
	ATOM	2597		LEU	179			114.255	1.00 31.68	В	Ċ
	ATOM	2598		LEU	179			2 114.747	1.00 31.31	В	C
55	ATOM	2599	C	LEU	179			3 113.425	1.00 31.91	В	Č
	ATOM	2600	ō	LEU	179			3 112.923	1.00 30.72	В	ō
	ATOM	2601	N	PRO	180			3 113.844	1.00 32.73	В	N
	ATOM	2602	CD	PRO	180			114.565	1.00 32.93	В	C
											_

	MOTA	2603	CA	PRO	180	6.932 -98.758 113.700 1.00 33.57 B	С
	ATOM	2604	СВ	PRO	180		C
	ATOM	2605	CG	PRO	180		C
	MOTA	2606	C	PRO	180		C
5		2607	0	PRO	180		
9	ATOM					· · · · · · · · · · · · · · · · · · ·	0
	ATOM	2608	N	VAL	181		N
	ATOM	2609	CA	VAL	181		С
	MOTA	2610	СВ	VAL	181		С
40	MOTA	2611		VAL	181		С
10	MOTA	2612		VAL	181		С
	MOTA	2613	С	VAL	181		С
	MOTA	2614	0	VAL	181		0
	MOTA	2615	N	PHE	182	3.599 ~96.079 114.538 1.00 35.49 B	N
	MOTA	2616	CA	PHE	182	2.726 -95.330 113.630 1.00 35.30 B	С
15	MOTA	2617	CB	PHE	182	3.431 -94.071 113.117 1.00 33.92 B	С
	MOTA	2618	CG	PHE	182		С
	MOTA	2619	CD1	PHE	182		С
	ATOM	2620	CD2	PHE	182		С
	ATOM	2621	CE1	PHE	182		С
20	ATOM	2622		PHE	182		Ċ
	ATOM	2623	CZ	PHE	182		C
	MOTA	2624	C	PHE	182		č
	ATOM	2625	ō	PHE	182		o
	ATOM	2626	N	ARG	183		N
25	ATOM	2627	CA	ARG	183		C
20	ATOM	2628	CB	ARG	183		C
	MOTA	2629	CG	ARG	183		C
	ATOM	2630	CD	ARG	183		
							C
30	MOTA	2631	NE	ARG	183		N
30	ATOM	2632	CZ	ARG	183		С
	MOTA	2633		ARG	183		N
	MOTA	2634		ARG	183		N
	MOTA	2635	C	ARG	183		С
25	MOTA	2636	0	ARG	183		0
35	MOTA	2637	N	SER	184		N
	ATOM	2638	CA	SER	184		С
	MOTA	2639	СВ	SER	184		С
	MOTA	2640	OG	SER	184		0
4.0	MOTA	2641	C	SER	184		С
40	MOTA	2642	0	SER	184		0
	ATOM	2643	N	LEU	185		N
	MOTA	2644	CA	LEU	185		C
	ATOM	2645	CB	LEU	185	-2.441 -97.131 113.330 1.00 43.53 B	С
	MOTA	2646	CG	LEU	185	-1.421 -96.007 113.534 1.00 44.02 B	С
45	ATOM	2647	CD1	LEU	185		С
	ATOM	2648	CD2	LEU	185	-0.417 -96.404 114.589 1.00 44.56 B	С
	ATOM	2649	С	LEU	185	-2.831 -98.510 111.271 1.00 42.68 B	С
	MOTA	2650	0	LEU	185		0
	ATOM	2651	N	PRO	186		N
50	MOTA	2652	CD	PRO	186	-5.129 -98.602 112.327 1.00 42.91 B	C
	MOTA	2653	CA	PRO	186	-4.877 -98.615 109.926 1.00 43.29 B	C
	ATOM	2654	СВ	PRO	186	-6.351 -98.617 110.337 1.00 43.26 B	c
	MOTA	2655	CG	PRO	186	-6.338 -97.972 111.696 1.00 43.70 B	Ċ
	ATOM	2656	c	PRO	186	-4.512 -97.421 109.054 1.00 43.49 B	C
55	ATOM	2657	Ö	PRO	186	-4.262 -96.326 109.556 1.00 43.14 B	o
-	ATOM	2658	N ·		187	-4.481 -97.650 107.747 1.00 43.61 B	И
	ATOM	2659	CA	ILE	187	-4.137 -96.613 106.784 1.00 43.56 B	C
		2660	CB	ILE	187		
	MOTA	2000	CB	TUC	10/	-4.406 -97.082 105.337 1.00 43.82 B	C

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	MOTA	2661	CG2		187		-96.018		1.00 43.79	В	С
	MOTA	2662		ILE	187		-98.395		1.00 44.06	В	С
	MOTA	2663		ILE	187		-98.288		1.00 44.66	В	С
_	ATOM	2664	С	ILE	187		-95.325		1.00 43.43	В	С
5	MOTA	2665	0	ILE	187		-94.232		1.00 42.67	В	0
	MOTA	2666	N	GLU	188		-95.451		1.00 43.55	В	N
	MOTA	2667	CA	GLU	188		-94.273		1.00 44.32	В	С
	MOTA	2668	CB	GLU	188		-94.658		1.00 45.62	В	С
4.0	MOTA	2669	CG	GLU	188			107.496	1.00 48.84	В	С
10	MOTA	2670	CD	GLU	188			106.406	1.00 50.12	В	С
	MOTA	2671	OE1		188			105.223	1.00 51.29	В	0
	MOTA	2672	OE2		188			106.737	1.00 50.87	В	0
	MOTA	2673	С	GLU	188			108.735	1.00 43.37	В	С
	MOTA	2674	0	GLU	188			108.748	1.00 43.27	В	0
15	ATOM	2675	N	ASP	189			109.776	1.00 42.33	В	N
	MOTA	2676	CA	ASP	189			110.979	1.00 41.76	В	С
	MOTA	2677	CB	ASP	189			112.134	1.00 42.89	В	С
	ATOM	2678	CG	ASP	189			112.806	1.00 44.75	В	С
00	MOTA	2679		ASP	189			113.879	1.00 46.20	В	0
20	MOTA	2680		ASP	189			112.258	1.00 45.42	В	0
	MOTA	2681	С	ASP	189			110.683	1.00 40.38	В	C
	MOTA	2682	0	ASP	189			111.102	1.00 40.12	В	0
	MOTA	2683	N	GLN	. 190			109.955	1.00 39.17	В	N
05	MOTA	2684	CA	GLN	190			109.592	1.00 38.44	В	С
25	MOTA	2685	CB	GLN	190			108.725	1.00 37.85	В	С
	MOTA	2686	CG	GLN	190			109.373	1.00 38.72	В	С
	MOTA	2687	CD	GLN	190			108.560	1.00 38.85	В	С
	MOTA	2688		GLN	190			108.789	1.00 38.95	В	0
20	MOTA	2689	NE2		190			107.611	1.00 39.18	В	N
30	MOTA	2690	C	GLN	190			108.822	1.00 37.77	В	C
	MOTA	2691	0	GLN	190			109.077	1.00 38.02	В	0
	ATOM	2692	N	ILE	191			107.877	1.00.37.50	В	N
	ATOM	2693	CA	ILE	191			107.061	1.00 37.36	В	С
25	MOTA	2694	CB	ILE	191			105.964	1.00 38.55	В	C
35	MOTA	2695	CG2		191			105.059	1.00 38.60	В	С
	MOTA	2696	CG1		191			105.138	1.00 39.51	В	С
	ATOM	2697	CD1		191			104.353	1.00 41.55	В	C
	ATOM	2698	C	ILE	191			107.919	1.00 36.51	В	C
40	ATOM	2699	0	ILE	191			107.804	1.00 36.70	В	0
40	ATOM	2700	N	SER	192			108.776	1.00 35.03	В	N
	ATOM	2701	CA	SER	192			109.646	1.00 34.26	В	С
	MOTA	2702	CB	SER	192			110.446	1.00 34.74		С
	ATOM	2703	OG	SER	192			109.586	1.00 35.76	В	0
45	MOTA	2704	C	SER	192			110.608	1.00 32.87	В	С
43	ATOM	2705	0	SER	192			110.853	1.00 31.92	В	0
	ATOM	2706	N	LEU	193			111.160	1.00 31.50	В	N
	MOTA	2707	CA	LEU	193			112.085	1.00 30.70	В	С
	ATOM	2708 2709	CB CG	LEU	193 193			112.786	1.00 29.93	В	C
50	ATOM			LEU				113.746	1.00 29.73	В	C
30	ATOM	2710		LEU	193			114.450	1.00 29.19	В	C
	MOTA	2711 2712	CD2	LEU	193			114.765	1.00 27.88	В	С
	MOTA	2712		LEU LEU	193			111.337	1.00 30.76	В	C
	ATOM	2713	O N		193			111.789	1.00 29.84	В	0
55	MOTA	2714	CA	LEU LEU	194 194			110.187	1.00 31.33	В	N
JJ	MOTA ATOM	2715	CB		194 194			109.389	1.00 32.09	В	С
		2716	CG	LEU	194 194			108.142	1.00 32.64	В	C
	MOTA	2717		LEU	194			107.333	1.00 34.23	В	C
	MOTA	2118	CDI	LEU	194	0.5/9	~00.995	106.557	1.00 35.50	В	С

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	MOTA	2719	CD2	LEU	194	2.116	-87.032	108.260	1.00 33.53	В	С
	ATOM	2720	С	LEU	194		-85.589		1.00 32.06	В	Č
	ATOM	2721	ō	LEU	194		-84.523		1.00 32.00		
	ATOM	2722	N	LYS	195		-85.651			В	0
5		2723							1.00 32.21	В	N
5	MOTA		CA	LYS	195		-84.448		1.00 32.26	В	C
	MOTA	2724	CB	LYS	195		-84.832		1.00 34.49	В	С
	MOTA	2725	CG	LYS	195		-85.427		1.00 36.99	В	C
	MOTA	2726	CD	LYS	195	-5.851	-86.146	105.619	1.00 38.55	В	С
	MOTA	2727	CE	LYS	195	-7.061	-85.223	105.632	1.00 39.77	В	С
10	MOTA	2728	NZ	LYS	195	-8.275	-85.939	105.131	1.00 41.50	В	N
	ATOM	2729	С	LYS	195	-3.541	-83.517	109.197	1.00 31.45	В	C
	MOTA	2730	0	LYS	195		-82.292		1.00 30.78	В	ō
	ATOM	2731	Ŋ	GLY	196		-84.105		1.00 29.85		
	ATOM	2732	CA	GLY	196		-83.303			В	N
15	MOTA	2733							1.00 28.37	В	С
13			С	GLY	196		-82.676		1.00 26.84	В	C
	MOTA	2734	0	GLY	196		-81.621		1.00 26.88	В	0
	MOTA	2735	N	ALA	197		-83.284		1.00 25.28	В	N
	MOTA	2736	CA	ALA	197		-82.751		1.00 23.92	В	С
	ATOM	2737	CB	ALA	197	-0.125	-83.776	113.816	1.00 23.40	В	С
20	ATOM	2738	С	ALA	197	0.608	-82.269	111.978	1.00 22.01	В	С
	ATOM	2739	0	ALA	197			112.550	1.00 20.56	В	ō
	ATOM	2740	N	ALA	198			110.668	1.00 20.97	В	N
	ATOM	2741	CA	ALA	198			109.836	1.00 20.46		C
	ATOM	2742	CB	ALA	198			108.349	1.00 20.48	В	
25	ATOM	2743	C	ALA	198			110.071		В	C
20									1.00 19.62	В	С
	ATOM	2744	0	ALA	198			110.365	1.00 19.14	В	0
	ATOM	2745	N	VAL	199			109.935	1.00 19.02	В	N
	MOTA	2746	CA	VAL	199			110.125	1.00 19.01	В	C
	MOTA	2747	CB	VAL	199			109.738	1.00 19.49	В	С
30	MOTA	2748		VAL	199	0.658	-75.938	110.073	1.00 18.87	В	С
	MOTA	2749	CG2	VAL	199	0.074	-77.563	108.243	1.00 19.44	В	С
	MOTA	2750	С	VAL	199			111.555	1.00 18.82	В	C.
	ATOM	2751	0	VAL	199			111.766	1.00 19.08	В	ō
	ATOM	2752	N	GLU	200			112.531	1.00 17.97	В	N
35	ATOM	2753	CA	GLU	200			113.934			
•	ATOM	2754	CB	GLU	200		-79.152		1.00 18.43	В	C
	ATOM	2755	CG						1.00 18.15	В	C
				GLU	200			115.030	1.00 20.07	В	С
	ATOM	2756	CD	GLU	200			115.866	1.00 21.56	В	С
40	MOTA	2757		GLU	200			116.765	1.00 21.43	В	0
40	MOTA	2758	OE2		200			115.630	1.00 22.35	В	0
	MOTA	2759	С	GLU	200	3.138	-78.854	114.202	1.00 17.94	В	С
	MOTA	2760	0	GLU	200	3.920	-78.148	114.830	1.00 18.17	В	0
	ATOM	2761	N	ILE	201	3.469	-80.047	113.716	1.00 17.97	В	N
	MOTA	2762	CA	ILE	201			113.897	1.00 17.91	В	C
45	MOTA	2763	CB	ILE	201			113.335	1.00 18.26	В	Č
. •	ATOM	2764	CG2		201			113.353	1.00 13.20	В	C
	ATOM	2765		ILE	201			114.233			C
		2766							1.00 17.28	В	C
	ATOM			ILE	201			113.701	1.00 18.61	В	C
50	ATOM	2767	С	ILE	201			113.255	1.00 17.32	В	C
50	ATOM	2768	0	ILE	201			113.815	1.00 18.72	В	0
	MOTA	2769	N	CYS	202			112.095	1.00 16.54	В	N
	ATOM	2770	CA	CYS	202			111.434	1.00 16.29	В	С
	MOTA	2771	CB	CYS	202	5.996	-77.845	110.062	1.00 16.39	В	С
	MOTA	2772	SG	CYS	202	5.924	-79.183	108.838	1.00 16.45	В	S
55	MOTA	2773	С	CYS	202	6.822	-77.049	112.295	1.00 15.32	В	C
	MOTA	2774	0	CYS	202			112.386	1.00 13.32	В	o
	ATOM	2775	Ŋ	HIS	203			112.924	1.00 14.13		
	ATOM	2776	CA	HIS	203			112.924		В	N
	AION	2,,0	CA	1113	203	0.061	-13.290	113./68	1.00 15.80	В	С

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	» mov	2777	СВ	HIS	203	4.750 -74.610 114.164 1.00 15.84 B	C
	MOTA MOTA	2778	CG	HIS	203	4.750 -74.610 114.164 1.00 15.84 B 4.173 -73.757 113.076 1.00 15.00 B	
	ATOM	2779	CD2		203	3.148 -73.977 112.221 1.00 15.47 B	_
	ATOM	2780	ND1		203	4.719 -72.545 112.714 1.00 15.56 B	
5	ATOM	2781	CE1		203	4.058 -72.056 111.681 1.00 15.06 B	
•	ATOM	2782	NE2	HIS	203	3.100 -72.906 111.361 1.00 15.27 B	
	ATOM	2783	C	HIS	203	6.886 -75.646 115.003 1.00 15.74 B	
	MOTA	2784	Ö	HIS	203	7.738 -74.867 115.422 1.00 16.61 B	
	MOTA	2785	N	ILE	204	6.643 -76.818 115.581 1.00 14.51 E	
10	ATOM	2786	CA	ILE	204	7.421 -77.263 116.734 1.00 15.77 E	
.0	MOTA	2787	СВ	ILE	204	6.914 -78.643 117.266 1.00 14.78 E	
	ATOM	2788		ILE	204	7.885 -79.192 118.305 1.00 14.01 E	
	ATOM	2789	CG1		204	5.496 -78.495 117.853 1.00 14.35 E	
	ATOM	2790		ILE	204	4.849 -79.812 118.317 1.00 11.99 E	
15	ATOM	2791	C	ILE	204	8.890 -77.404 116.303 1.00 15.99 E	
	ATOM	2792	Ö	ILE	204	9.803 -76.963 117.001 1.00 14.96 E	
	ATOM	2793	N	VAL	205	9.108 -78.011 115.140 1.00 16.33 E	
	ATOM	2794	CA	VAL	205	10.463 -78.208 114.623 1.00 18.21 E	
	ATOM	2795	CB	VAL	205	10.455 -79.139 113.402 1.00 17.84 E	
20	ATOM	2796		VAL	205	11.796 -79.058 112.671 1.00 17.90 E	
	ATOM	2797		VAL	205	10.169 -80.565 113.862 1.00 18.55 E	
	ATOM	2798	C	VAL	205	11.153 -76.906 114.242 1.00 17.83 F	
	ATOM	2799	0	VAL	205		3 0
	ATOM	2800	N	LEU	206		3 N
25	MOTA	2801	CA	LEU	206		3 C
	ATOM	2802	CB	LEU	206		3 C
	ATOM	2803	CG	LEU	206		3 C
	ATOM	2804		LEU	206		3 C
	ATOM	2805		LEU	206		3 C
30	ATOM	2806	С	LEU	206		в с
	MOTA	2807	0	LEU	206		в о
	ATOM	2808	N	ASN	207		в и
	ATOM	2809	CA	ASN	207	10.918 -73.251 116.680 1.00 17.40	в С
	MOTA	2810	CB	ASN	207	10.124 -73.774 117.874 1.00 15.99	в с
35	MOTA	2811	CG	ASN	207	10.184 -72.828 119.073 1.00 16.60	в с
	MOTA	2812	OD1	ASN	207	10.860 -73.101 120.068 1.00 15.13	в о
	MOTA	2813	ND2	ASN	207	9.485 -71.701 118.970 1.00 13.67	B N
	ATOM	2814	С	ASN	207	12.387 -73.144 117.061 1.00 18.07	в с
	ATOM	2815	0	ASN	207	12.804 -72.133 117.621 1.00 17.57	в о
40	ATOM	2816	N	THR	208		B N
	MOTA	2817	CA	THR	208	14.595 -74.147 117.080 1.00 20.72	в с
	ATOM	2818	CB	THR	208		в с
	ATOM	2819		THR	208		в о
	MOTA	2820	CG2	THR	208		в с
45	MOTA	2821	С	THR	208		в с
	ATOM	2822	0	THR	208		в о
	MOTA	2823	N	THR	209		в и
	ATOM	2824	CA	THR	209		в с
	ATOM	2825	CB	THR	209		в с
50	MOTA	2826	OG1		209		в о
	MOTA	2827	CG2		209		в с
	ATOM	2828	С	THR	209		в с
	MOTA	2829	0	THR	209		в о
	MOTA	2830	N	PHE	210		B N
55	MOTA	2831	CA	PHE	210		в с
	MOTA	2832	СВ	PHE	210		в с
	MOTA	2833	CG	PHE	210		в с
	MOTA	2834	CD1	PHE	210	10.694 -67.081 115.914 1.00 18.35	в с

	3 mov	2025	CD2	DUE	210	11 161	-67.046	110 262	1 00 10 05	-	_
	MOTA	2835			210			116.203	1.00 19.05	В	C
	MOTA	2836	CE1						1.00 18.54	В	C
	MOTA	2837	CE2	PHE	210		-65.865		1.00 19.42	В	C
5	ATOM	2838	CZ	PHE	210			117.286	1.00 18.58	В	C
5	MOTA	2839	С	PHE	210			116.986	1.00 20.77	В	C
	MOTA	2840	0	PHE	210			118.021	1.00 20.70	В	0
	MOTA	2841	N	CYS	211			116.565	1.00 22.21	В	N
	MOTA	2842	CA	CYS	211			117.327	1.00 24.26	В	C
	MOTA	2843	СВ	CYS	211			116.391	1.00 24.98	В	С
10	MOTA	2844	SG	CYS	211			117.267	1.00 27.94	В	S
	MOTA	2845	С	CYS	211			118.148	1.00 25.17	В	C
	MOTA	2846	0	CYS	211			117.586	1.00 24.50	В	0
	MOTA	2847	N	LEU	212	14.950	-65.076	119.471	1.00 25.78	В	N
	MOTA	2848	CA	LEU	212	14.173	-64.229	120.375	1.00 27.77	В	С
15	MOTA	2849	CB	LEU	212	14.396	-64.652	121.830	1.00 26.74	В	C
	MOTA	2850	CG	LEU	212	13.735	-65.968	122.237	1.00 26.59	В	С
	MOTA	2851	CD1	LEU	212	14.046	-66.279	123.700	1.00 26.25	В	С
	MOTA	2852	CD2	LEU	212	12.235	-65.863	122.014	1.00 24.89	В	С
	ATOM	2853	С	LEU	212	14.448	-62.739	120.253	1.00 28.72	В	С
20	ATOM	2854	0	LEU	212	13.521	-61.931	120.271	1.00 29.38	В	0
	ATOM	2855	N	GLN	213	15.720	-62.379	120.140	1.00 30.42	В	N
	MOTA	2856	CA	GLN	213			120.027	1.00 32.34	В	C
	MOTA	2857	СB	GLN	213	17.631	-60.869	119.893	1.00 35.12	В	С
	MOTA	2858	CG	GLN	213	18.122	-59.462	119.576	1.00 38.96	В	C
25	MOTA	2859	CD	GLN	213	18.458	-58.655	120.817	1.00 41.53	В	C
	ATOM	2860	OE1	GLN	213	19.566	-58.754	121.357	1.00 42.98	В	0
	ATOM	2861	NE2	GLN	213	17.500	-57.858	121.284	1.00 42.35	В	N
	ATOM	2862	С	GLN	213	15.453	-60.262	118.846	1.00 32.03	В	С
	MOTA	2863	0	GLN	213	15.005	-59.126	118.986	1.00 32.80	В	0
30	ATOM	2864	N	THR	214			117.691	1.00 30.84	В	N
	MOTA	2865	CA	THR	214	14.828	-60.288	116.497	1.00 29.82	В	С
	ATOM	2866	CB	THR	214			115.316	1.00 29.30	В	C
	MOTA	2867	OG1	THR	214			115.049	1.00 28.95	В	0
	MOTA	2868	CG2		214	17.108	-59.679	115.636	1.00 30.14	В	C
35	ATOM	2869	С	THR	214	13.472	-60.792	116.009	1.00 29.36	В	С
	MOTA	2870	0	THR	214			115.032	1.00 28.67	В	0
	MOTA	2871	N	GLN	215			116.670	1.00 28.97	В	N
	ATOM	2872	CA	GLN	215			116.273	1.00 29.50	В	С
	ATOM	2873	СВ	GLN	215			116.375	1.00 30.88	В	Ċ
40	MOTA	2874	CG	GLN	215			117.764	1.00 33.91	В	C
	ATOM	2875	CD	GLN	215			118.726	1.00 35.71	В	Ċ
	MOTA	2876		GLN	215			118.513	1.00 37.61	В	ō
	ATOM	2877		GLN	215			119.793	1.00 36.76	В	N
	ATOM	2878	С	GLN	215			114.834	1.00 28.46	В	C
45	MOTA	2879	0	GLN	215			114.093	1.00 28.52	В	ō
	ATOM	2880	N	ASN	216			114.442	1.00 26.98	В	N
	ATOM	2881	CA	ASN	216			113.092	1.00 25.74	В	C
	ATOM	2882	СВ	ASN	216			112.363	1.00 27.52	В	c
	ATOM	2883	CG	ASN	216			112.093	1.00 28.70	В	Č
50	ATOM	2884		ASN	216			111.749	1.00 27.69	В	ŏ
	ATOM	2885		ASN	216			112.228	1.00 29.92	В	N
	ATOM	2886	C	ASN	216			113.125	1.00 24.08	В	C
	MOTA	2887	Ö	ASN	216			114.119	1.00 22.97	В	ŏ
	ATOM	2888	Ŋ	PHE	217			112.032	1.00 22.85	В	N
55	ATOM	2889	CA	PHE	217			111.906	1.00 22.30	В	C
-	MOTA	2890	СВ	PHE	217			111.195	1.00 20.84	В	c
	MOTA	2891	CG	PHE	217			111.975	1.00 20.02	В	
	ATOM	2892		PHE	217			111.948	1.00 20.02	В	
						_0.000					_

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	ATOM	2893	CD2	PHE	217	10.805	-69.644	112.733	1.00 19.92	В	С
	MOTA	2894	CE1	PHE	217		-67.622		1.00 18.54	В	С
	MOTA	2895	CE2	PHE	217	9.620	-69.780	113.453	1.00 19.66	В	C
_	MOTA	2896	CZ	PHE	217	8.672	-68.763	113.415	1.00 19.48	В	С
5	ATOM	2897	C	PHE	217		-67.588		1.00 22.98	В	C
	ATOM	2898	0	PHE	217	14.652	-67.184	109.895	1.00 21.70	В	0
	ATOM	2899	N	LEU	218	15.777	-68.046	111.645	1.00 23.53	В	N
	MOTA	2900	CA	LEU	218	17.066	-68.114	110.957	1.00 24.87	В	C
	MOTA	2901	CB	LEU	218	18.210	-67.785	111.930	1.00 26.08	В	C
10	MOTA	2902	CG	LEU	218	18.380	-66.339	112.436	1.00 27.99	В	C
	MOTA	2903	CD1		218		-65.845		1.00 27.86	В	С
	ATOM	2904	CD2	LEU	218	19.555	-66.279	113.413	1.00 27.96	В	С
	MOTA	2905	С	LEU	218	17.260	-69.510	110.383	1.00 25.07	В	C
	MOTA	2906	0	LEU	218	17.539	-70.458	111.115	1.00 25.64	В	0
15	ATOM	2907	N	CYS	219	17.105	-69.631	109.069	1.00 24.23	В	N
	ATOM	2908	CA	CYS	219	17.244	-70.918	108.396	1.00 23.85	В	С
	ATOM	2909	CB	CYS	219	15:935	-71.263	107.674	1.00 22.48	В	C
	MOTA	2910	SG	CYS	219	14.462	-71.225	108.747	1.00 19.92	В	s
	MOTA	2911	С	CYS	219	18.406	-70.868	107.398	1.00 24.42	В	С
20	ATOM	2912	0	CYS	219	18.231	-70.505	106.227	1.00 22.82	В	0
	MOTA	2913	N	GLY	220	19.590	-71.252	107.870	1.00 25.23	В	N
	MOTA	2914	CA	GLY	220	20.764	-71.214	107.021	1.00 24.90	В	С
	ATOM	2915	С	GLY	220	20.996	-69.754	106.695	1.00 24.77	В	C
	ATOM	2916	0	GLY	220		-68.922		1.00 24.69	В	0
25	MOTA	2917	N	PRO	221		-69.401		1.00 24.50	В	N
	ATOM	2918	CD	PRO	221		-70.265		1.00 23.93	В	C
	ATOM	2919	CA	PRO	221		-67.993		1.00 24.42	В	C
	ATOM	2920	СВ	PRO	221			103.699	1.00 23.72	В	C
	ATOM	2921	CG	PRO	221			103.085	1.00 23.75	В	C
30	ATOM	2922	С	PRO	221	20.073	-67.181	104.995	1.00 23.85	В	C
	ATOM	2923	0	PRO	221			104.811	1.00 23.61	В	0
	ATOM	2924	N	LEU	222			105.153	1.00 22.94	В	N
	MOTA	2925	CA	LEU	222			105.084	1.00 22.55	В	C
	ATOM	2926	СВ	LEU	222			104.478	1.00 20.79	В	C
35	ATOM	2927	CG	LEU	222			103.100	1.00 19.76	В	C
	ATOM	2928		LEU	222			102.623	1.00 20.37	В	C
	ATOM	2929		LEU	222			102.122	1.00 20.15	В	C
	ATOM	2930	С	LEU	222			106.415	1.00 22.84	В	C
	ATOM	2931	0	LEU	222			107.473	1.00 23.24	В	O
40	MOTA	2932	N	ARG	223			106.333	1.00 22.89	В	N
	ATOM	2933	CA	ARG	223			107.485	1.00 24.01	В	C
	ATOM	2934	СВ	ARG	223	16.451	-63.650	107.782	1.00 26.47	В	C
	ATOM	2935	CG	ARG	223			108.768	1.00 31.77	В	С
	ATOM	2936	CD	ARG	223			108.616	1.00 35.27	В	C
45	ATOM	2937	NE	ARG	223			107.349	1.00 38.31	В	N
	ATOM	2938	cz	ARG	223	16.314	-59.714	106.740	1.00 39.39	В	C
	ATOM	2939		ARG	223			107.279	1.00 41.22	В	N
	ATOM	2940		ARG	223			105.600	1.00 40.79	В	N
	ATOM	2941	C	ARG	223			107.183	1.00 23.59	В	C
50	ATOM	2942	0	ARG	223			106.287	1.00 22.38	В	0
•••	ATOM	2943	N	TYR	224			107.930	1.00 22.01	В	N
	ATOM	2944	CA	TYR	224			107.742	1.00 21.48	В	C
	MOTA	2945	СВ	TYR	224			107.718	1.00 20.04	В	Ċ
	MOTA	2946	CG	TYR	224			106.615	1.00 19.13	В	C
55	ATOM	2947		TYR	224			106.851	1.00 17.91	В	C
-	ATOM	2948		TYR	224			105.848	1.00 17.51	В	C
	MOTA	2949		TYR	224			105.338	1.00 17.45	В	C
	ATOM	2950		TYR	224			104.319	1.00 17.93	В	C
	N 1 OLI	2750			227	552	JJ.123		1.00 17.93		C

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	MOTA	2951	CZ	TYR	224	12.328	-69.118	104.587	1.00 17.71	В	С
	MOTA	2952	OH	TYR	224	12.714	-69.995	103.606	1.00 17.71	В	0
	MOTA	2953	С	TYR	224	11.352	-64.239	108.872	1.00 21.91	В	С
_	MOTA	2954	0	TYR	224			110.043	1.00 21.72	В	0
5	MOTA	2955	N	THR	225			108.509	1.00 21.54	В	N
	ATOM	2956	CA	THR	225			109.478	1.00 21.65	В	С
	MOTA	2957	CB	THR	225	10.335	-60.874	109.202	1.00 23.19	В	C
	MOTA	2958		THR	225			107.912	1.00 22.96	В	0
	MOTA	2959	CG2	THR	225			109.232	1.00 23.33	В	С
10	ATOM	2960	С	THR	225			109.416	1.00 20.84	В	С
	MOTA	2961	0	THR	225			108.513	1.00 19.99	В	0
	MOTA	2962	N	ILE	226			110.376	1.00 19.42	В	N
	MOTA	2963	CA	ILE	226			110.430	1.00 18.00	В	C
	MOTA	2964	CB	ILE	226			111.768	1.00 17.71	В	С
15	MOTA	2965	CG2	ILE	226			111.813	1.00 16.28	В	С
	MOTA	2966	CG1		226			111.963	1.00 15.92	В	С
	MOTA	2967	CD1	ILE	226			113.363	1.00 13.01	В	С
	MOTA	2968	С	ILE	226			109.207	1.00 18.53	В	С
~~	MOTA	2969	0	ILE	226			108.754	1.00 17.76	В	0
20	MOTA	2970	N	GLU	227			108.649	1.00 18.32	В	N
	MOTA	2971	CA	GLU	227			107.468	1.00 19.11	В	C
	MOTA	2972	СВ	GLU	227			107.103	1.00 20.59	В	C
	ATOM	2973	CG	GLU	227			107.959	1.00 20.58	В	С
0.5	MOTA	2974	CD	GLU	227		-57.143		1.00 21.61	В	С
25	ATOM	2975		GLU	227			109.617	1.00 23.10	В	0
	MOTA	2976		GLU	227			110.308	1.00 22.37	В	0
	MOTA	2977	C	GLU	227			106.273	1.00 18.86	В	С
	MOTA	2978	0	GLU	227			105.416	1.00 18.23	В	0
00	ATOM	2979	N	ASP	228			106.199	1.00 18.37	В	N
30	MOTA	2980	CA	ASP	228			105.071	1.00 18.69	В	С
	MOTA	2981	CB	ASP	228			105.088	1.00 18.93	В	С
	MOTA	2982	CG	ASP	228			104.950	1.00 19.98	В	С
	ATOM	2983		ASP	228			104.031	1.00 20.95	В	0
25	ATOM	2984		ASP	228			105.754	1.00 19.87	В	0
35	MOTA	2985	С	ASP	228			105.102	1.00 18.01	В	С
	ATOM	2986	0	ASP	228			104.064	1.00 18.44	В	0
	MOTA	2987	N	GLY	229			106.289	1.00 17.20	В	N
	MOTA	2988	CA	GLY	229			106.392	1.00 17.52	В	С
40	ATOM	2989	С	GLY	229			106.035	1.00 17.70	В	C
40	ATOM	2990	0	GLY	229			105.330	1.00 18.08	В	0
	ATOM	2991	N	ALA	230			106.510	1.00 17.31	В	N
	ATOM	2992	CA	ALA	230			106.237		В	
	ATOM	2993	СВ	ALA	230			107.031	1.00 17.60	В	C
45	ATOM	2994	C	ALA	230			104.749	1.00 18.51	В	C
45	MOTA	2995	0	ALA	230			104.231	1.00 17.21	В	0
	MOTA	2996	N	ARG	231 231			104.060	1.00 19.86	В	N
	MOTA	2997	CA	ARG				102.643	1.00 20.24	В	C
	MOTA	2998 2999	CB CG	ARG	231 231			102.173 102.843	1.00 21.54	В	C
50	ATOM			ARG					1.00 23.12	В	C
50	MOTA	3000	CD	ARG	231			102.506 101.092	1.00 23.94	В	C
	ATOM	3001 3002	NE CZ	ARG	231				1.00 25.20	В	И
	ATOM	3002		ARG ARG	231 231			100.553 101.301	1.00 25.86	В	C
	ATOM	3003					-55.746		1.00 25.92	В	N
55	MOTA	3004	C	ARG ARG	231 231				1.00 26.68	В	N
JJ	MOTA MOTA	3005	0	ARG	231			101.765 100.640	1.00 20.12	В	C
		3006	Ŋ	VAL	231			100.640	1.00 19.52	В	0
	MOTA		CA	VAL	232			102.256	1.00 19.90	В	N
	ATOM	3008	CA	VAL	434	1.239	-03.195	101.450	1.00 19.34	В	С

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	MOTA	3009		VAL	232	1.927 -64.592 101.612 1.00 20.66 B	С
	MOTA	3010	CG1		232	3.413 -64.476 101.307 1.00 19.33 B	С
	MOTA	3011	CG2		232	1.671 -65.169 103.017 1.00 18.79 B	С
_	MOTA	3012	С	VAL	232	-0.245 -63.330 101.779 1.00 19.07 B	С
5	MOTA	3013	0	VAL	232	-0.926 -64.191 101.230 1.00 17.55 B	0
	MOTA	3014	N	GLY	233	-0.745 -62.486 102.680 1.00 19.78 B	N
	MOTA	3015	CA	GLY	233	-2.161 -62.535 103.008 1.00 20.41 B	С
	MOTA	3016	C	GLY	233	-2.643 -62.915 104.397 1.00 21.13 B	C
40	MOTA	3017	0	GLY	233	-3.805 -62.655 104.719 1.00 21.49 B	0
10	MOTA	3018	N	PHE	234	-1.806 -63.541 105.223 1.00 20.51 B	N
	MOTA	3019	CA	PHE	234	-2.268 -63.905 106.563 1.00 20.56 B	С
	ATOM	3020	СВ	PHE	234	-1.205 -64.706 107.326 1.00 20.73 B	С
	MOTA	3021	CG	PHE	234	-0.998 -66.103 106.810 1.00 20.24 B	C
A.E.	MOTA	3022	CD1		234	0.169 -66.441 106.135 1.00 19.02 B	С
15	MOTA	3023	CD2		234	-1.956 -67.093 107.035 1.00 19.67 B	C
	MOTA	3024	CE1		234	0.386 -67.747 105.695 1.00 19.63 B	С
	MOTA	3025		PHE	234	-1.748 -68.403 106.597 1.00 19.47 B	С
	MOTA	3026	CZ	PHE	234	-0.574 -68.731 105.928 1.00 18.87 B	С
00	MOTA	3027	С	PHE	234	-2.602 -62.651 107.362 1.00 20.67 B	С
20	MOTA	3028	0	PHE	234	-1.936 -61.624 107.234 1.00 20.71 B	0
	MOTA	3029	N	GLN	235	-3.632 -62.740 108.194 1.00 21.02 B	N
	ATOM	3030	CA	GLN	235	-4.051 -61.614 109.020 1.00 22.09 B	С
	ATOM	3031	CB	GLN	235	-5.468 -61.845 109.548 1.00 22.57 B	С
05	MOTA	3032	CG	GLN	235	-6.481 -61.997 108.428 1.00 24.45 B	С
25	MOTA	3033	CD	GLN	235	-7.898 -62.104 108.926 1.00 27.02 B	C
	ATOM	3034		GLN	235	-8.219 -62.966 109.748 1.00 27.06 B	0
	MOTA	3035	NE2		235	-8.768 -61.229 108.423 1.00 27.49 B	N
	MOTA	3036	C	GLN	235	-3.085 -61.414 110.173 1.00 21.61 B	С
20	ATOM	3037	0	GLN	235	-2.546 -62.372 110.718 1.00 21.23 B	0
30	MOTA	3038	N .	VAL	236	-2.876 -60.158 110.542 1.00 21.74 B	N
	MOTA	3039	CA	VAL	236	-1.957 -59.813 111.616 1.00 22.27 B	С
	ATOM	3040	CB	VAL	236	-1.915 -58.278 111.810 1.00 22.19 B	C
	ATOM	3041		VAL	236	-1.086 -57.914 113.032 1.00 22.00 B	C
25	ATOM	3042		VAL	236	-1.329 -57.628 110.562 1.00 21.53 B	C
35	MOTA	3043	C	VAL	236	-2.250 -60.503 112.944 1.00 22.51 B	С
	ATOM	3044	0	VAL	236	-1.330 -60.979 113.604 1.00 22.18 B	0
	ATOM	3045	N	GLU	237	-3.521 -60.558 113.332 1.00 22.41 B	N
	ATOM	3046	CA	GLU	237	-3.928 -61.198 114.586 1.00 22.96 B	C
40	MOTA	3047	CB	GLU	237	-5.452 -61.107 114.748 1.00 25.41 B	C
40	ATOM ATOM	3048 3049	CG CD	GLU GLU	237 237	-6.012 -61.912 115.905 1.00 29.15 B -7.516 -61.721 116.076 1.00 31.86 B	C
	ATOM	3049		GLU	237		
					237		
	MOTA MOTA	3051 3052	C	GLU GLU	237		0
45	ATOM	3053	0	GLU	237		C
70	ATOM	3054	N	PHE	238		0
	ATOM	3055	CA	PHE	238		N
	ATOM	3056	CB	PHE	238		C
	ATOM	3057	CG	PHE	238	-3.746 -65.287 112.033 1.00 18.50 B -3.147 -66.608 111.679 1.00 18.24 B	C
50	ATOM	3058		PHE	238		
50	MOTA	3059		PHE	238		C
	ATOM	3060		PHE	238		C
	ATOM	3061		PHE	238		C
		3062	CEZ	PHE	238		C
55	ATOM ATOM	3062	C	PHE	238	-1.994 -69.072 111.025 1.00 19.34 B -1.777 -64.831 113.526 1.00 19.52 B	C
JJ	ATOM	3064	0	PHE	238		С
	ATOM	3065	N	LEU	239	-1.243 -65.653 114.276 1.00 17.63 B -1.087 -63.957 112.799 1.00 19.51 B	0
		3066	CA	LEU	239		N
	MOTA	2000	CA	TEO	233	0.372 -63.903 112.840 1.00 21.27 B	С

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	MOTA	3067		LEU	239		-62.786		1.00 19.88	В	С
	MOTA	3068		LEU	239		-63.090		1.00 21.98	В	C
	MOTA	3069	CD1		239	0.983	-64.488	110.062	1.00 19.74	В	С
_	MOTA	3070	CD2	LEU	239	0.798	-62.046	109.521	1.00 20.05	В	C
5	ATOM	3071	С	LEU	239	0.876	-63.678	114.268	1.00 21.93	В	С
	MOTA	3072	0	LEU	239	1.819	-64.347	114.711	1.00 21.20	В	0
	MOTA	3073	N	GLU	240	0.244	-62.752	114.991	1.00 22.27	В	N
	ATOM	3074	CA	GLU	240		-62.455		1.00 24.45	В	C
	ATOM	3075		GLU	240		-61.313		1.00 26.77	В	Č
10	ATOM	3076		GLU	240		-59.949		1.00 30.99	В	Ċ
	ATOM	3077		GLU	240		-59.403		1.00 34.68	В	Ċ
	ATOM	3078	OE1		240		-59.754		1.00 34.00	В	Ö
	ATOM	3079	OE2		240		-58.604		1.00 36.81		Ö
	ATOM	3080	C	GLU	240		-63.689		1.00 38.81	B B	c
15	ATOM	3081	Ö	GLU	240		-63.952				
13	ATOM	3082							1.00 23.70	В	0
			N	LEU	241			117.087	1.00 22.32	В	N
	MOTA	3083	CA	LEU	241			117.864	1.00 21.98	В	С
	MOTA	3084	CB	LEU	241			117.517	1.00 23.58	В	C
00	MOTA	3085	CG	LEU	241			118.172	1.00 25.77	В	С
20	MOTA	3086	CD1		241			119.674	1.00 26.71	В	С
	MOTA	3087	CD2		241			117.550	1.00 27.49	В	С
	MOTA	3088	С	LEU	241			117.544	1.00 20.98	В	C
	MOTA	3089	0	LEU	241	0.858	-67.281	118.434	1.00 20.13	В	0
	MOTA	3090	N	LEU	242	0.666	-66.746	116.265	1.00 19.66	В	N
25	ATOM	3091	CA	LEU	242	1.711	-67.663	115.827	1.00 19.39	В	С
	MOTA	3092	CB	LEU	242	1.757	-67.706	114.296	1.00 17.57	В	С
	MOTA	3093	CG	LEU	242	2.800	-68.626	113.656	1.00 17.99	В	С
	MOTA	3094	CD1	LEU	242			114.206	1.00 17.20	В	C
	ATOM	3095		LEU	242			112.148	1.00 17.85	В	Č
30	ATOM	3096	С	LEU	242			116.380	1.00 19.08	В	Č
	ATOM	3097	0	LEU	242			116.858	1.00 18.27	В	ō
	ATOM	3098	N	PHE	243			116.322	1.00 19.06	В	N
	ATOM	3099	CA	PHE	243			116.820	1.00 20.50	В	C
	ATOM	3100	СВ	PHE	243			116.020	1.00 20.80	В	C
35	MOTA	3101	CG	PHE	243			114.866	1.00 20.80	В	C
55	ATOM	3101		PHE	243			113.773			
	ATOM	3102		PHE	243				1.00 21.05	В	C
								114.665	1.00 20.91	В	C
	ATOM	3104		PHE	243			112.497	1.00 21.45	В	C
40	ATOM	3105		PHE	243			113.395	1.00 21.14	В	C
40	MOTA	3106	CZ	PHE	243			112.308	1.00 21.10	В	С
	MOTA	3107	C	PHE	243			118.340	1.00 21.66	В	С
	MOTA	3108	0	PHE	243			118.904	1.00 21.34	В	0
	MOTA	3109	N	HIS	244			118.995	1.00 22.31	В	N
	MOTA	3110	CA	HIS	244			120.448	1.00 23.28	В	C
45	MOTA	3111	СВ	HIS	244			120.951	1.00 26.24	В	C
	MOTA	3112	CG	HIS	244			122.391	1.00 29.43	В	C
	MOTA	3113	CD2	HIS	244	2.047	-64.644	123.516	1.00 30.00	В	С
	MOTA	3114	ND1	HIS	244	1.623	-66.666	122.804	1.00 30.79	В	N
	MOTA	3115	CE1	HIS	244	1.494	-66.682	124.119	1.00 30.26	В	C
50	MOTA	3116	NE2	HIS	244	1.747	-65.469	124.575	1.00 30.77	В	N
	MOTA	3117	С	HIS	244	3.904	-66.892	120.857	1.00 21.80	В	C
	ATOM	3118	0	HIS	244			121.809	1.00 21.67	В	ō
	ATOM	3119	N	PHE	245			120.137	1.00 19.90	В	Ŋ
	ATOM	3120	CA	PHE	245			120.389	1.00 18.67	В	c
55	ATOM	3121	CB	PHE	245			119.356	1.00 18.04	В	Č
	MOTA	3122	CG	PHE	245			119.316	1.00 17.45	В	c
	ATOM	3123		PHE	245			120.294	1.00 17.43	В	C
	MOTA	3124		PHE	245			118.321	1.00 10.43	В	C
	ATOM	7124	CDZ	ERE	243	4.00/	-/1.303	110.321	1.00 1/.18	D	C

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	MOTA	3125	CE1	PHE	245	3.244	-73.760	120.286	1.00 16.09	В	С
	MOTA	3126	CE2	PHE	245	4.539	-73.306	118.303	1.00 17.01	В	C
	MOTA	3127	CZ	PHE	245	4.117	-74.196	119.287	1.00 16.45	В	С
	MOTA	3128	С	PHE	245	5.030	-69.556	120.290	1.00 17.95	В	С
5	MOTA	3129	0	PHE	245	5.600	-70.169	121.183	1.00 17.52	В	0
	MOTA	3130		HIS	246	5.666	-69.115	119.206	1.00 16.92	В	N
	ATOM	3131		HIS	246			119.024	1.00 16.61	В	C
	ATOM	3132		HIS	246			117.620	1.00 15.88	В	Ċ
	ATOM	3133		HIS	246			116.554	1.00 16.90	В	c
10	ATOM	3134	CD2		246			115.599	1.00 17.28	В	C
.0	MOTA	3135	ND1		246			116.440	1.00 17.28	В	
	ATOM	3136	CE1		246			115.465			N
									1.00 16.06	В	C
	ATOM	3137	NE2		246			114.940	1.00 15.84	В	N
4 5	MOTA	3138	C	HIS	246			120.095	1.00 16.14	В	С
15	MOTA	3139	0	HIS	246			120.598	1.00 16.15	В	0
	MOTA	3140	N	GLY	247			120.445	1.00 15.88	В	N
	ATOM	3141	CA	GLY	' 247			121.488	1.00 16.58	В	С
	MOTA	3142	С	GLY	247	8.326	-67.552	122.820	1.00 17.29	В	С
	MOTA	3143	0	GLY	247	9.325	-67.849	123.474	1.00 16.50	В	0
20	MOTA	3144	N	THR	248	7.097	-67.872	123.215	1.00 17.62	В	N
	MOTA	3145	CA	THR	248	6.854	-68.569	124.473	1.00 19.52	В	С
	MOTA	3146	CB	THR	248	5.334	-68.737	124.715	1.00 20.24	В	С
	ATOM	3147		THR	248			124.589	1.00 19.56	В	O
	ATOM	3148	CG2	THR	248			126.110	1.00 20.09	В	Č
25	ATOM	3149	C	THR	248			124.530	1.00 19.85	В	C
	ATOM	3150	Õ	THR	248			125.519	1.00 18.96	В	ō
	ATOM	3151	N	LEU	249			123.466	1.00 10.90	В	И
	ATOM	3152	CA	LEU	249			123.412	1.00 13.33	В	
					249						C
20	ATOM	3153	CB	LEU				122.145	1.00 21.21	В	C
30	MOTA	3154	CG	LEU	249			121.875	1.00 19.48	В	C
	MOTA	3155		LEU	249			123.015	1.00 21.12	В	C
	MOTA	3156		LEU	249			120.547	1.00 19.38	В	С
	MOTA	3157	С	LEU	249			123.429	1.00 22.24	В	С
	MOTA	3158	0	LEU	249			124.101	1.00 21.93	В	0
35	MOTA	3159	N	ARG	250	10.058	-70.981	122.683	1.00 23.40	В	N
	MOTA	3160	CA	ARG	250			122.592	1.00 25.09	В	С
	MOTA	3161	СВ	ARG	250	11.789	-69.621	121.613	1.00 26.51	В	C
	ATOM	3162	CG	ARG	250	12.705	-69.985	120.465	1.00 28.87	В	C
	MOTA	3163	CD	ARG	250	14.055	-70.492	120.927	1.00 28.96	В	C
40	ATOM	3164	NE	ARG	250			119.817	1.00 29.63	В	N
	ATOM	3165	CZ	ARG	250	16.291	-70.825	119.918	1.00 31.64	В	С
	ATOM	3166	NH1	ARG	250			118.839	1.00 32.55	В	N
	ATOM	3167		ARG	250			121.087	1.00 32.97	В	N
	ATOM	3168	С	ARG	250			123.932	1.00 25.34	В	C
45	ATOM	3169	Ö	ARG	250			124.285	1.00 24.47	В	ō
.0	ATOM	3170	N	LYS	251			124.667	1.00 24.47	В	
	ATOM	3171	CA	LYS	251			125.963	1.00 20.27		N
		3172	CB		251			126.536		В	C
	ATOM			LYS					1.00 27.26	В	C
EΩ	ATOM	3173	CG	LYS	251			125.798	1.00 30.10	В	C
50	MOTA	3174	CD	LYS	251			126.269	1.00 32.49	В	С
	MOTA	3175	CE	LYS	251			125.419	1.00 34.43	В	С
	MOTA	3176	NZ	LYS	251			125.733	1.00 37.02	В	N
	MOTA	3177	С	LYS	251			126.982	1.00 26.91	В	С
	MOTA	3178	0	LYS				127.980	1.00 27.68	В	0
55	MOTA	3179	N	LEU				126.739	1.00 26.21	В	N
	MOTA	3180	CA	LEU		11.577	-72.462	2 127.648	1.00 25.54	В	С
	MOTA	3181	CB	LEU	252	10.405	-73.412	127.395	1.00 24.47	В	С
	ATOM	3182	CG	LEU	252			127.809	1.00 23.94	В	C
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	MOTA	3183	CD1		252		-73.892		1.00 21.51	В	С
	MOTA	3184	CD2	LEU	252	9.026	-72.598	129.317	1.00 22.42	В	С
	MOTA	3185	С	LEU	252	12.904	-73.224	127.552	1.00 26.17	В	С
	MOTA	3186	0	LEU	252	13.200	-74.077	128.399	1.00 25.06	В	0
5	ATOM	3187	N	GLN	253		-72.920		1.00 26.13	В	N
	ATOM	3188	CA	GLN	253		-73.571		1.00 27.67	В	C
	ATOM	3189	СВ	GLN	253		-73.124		1.00 29.23	В	C
	ATOM	3190	CG	GLN	253		-71.621		1.00 23.23	В	C
	MOTA	3191	CD	GLN	253 253			128.673			
10									1.00 34.60	В	C
10	MOTA	3192	OE1		253			128.552	1.00 35.64	В	0
	MOTA	3193	NE2	GLN	253			129.770	1.00 35.92	В	N
	MOTA	3194	C	GLN	253			126.392	1.00 27.27	В	С
	MOTA	3195	0	GLN	253			127.170	1.00 27.62	В	0
	ATOM	3196	N	LEU	254	13.979	-75.659	125.578	1.00 26.25	В	N
15	MOTA	3197	CA	LEU	254	13.781	-77.100	125.583	1.00 25.87	В	С
	MOTA	3198	CB	LEU	254	12.541	-77.484	124.763	1.00 23.25	В	С
	ATOM	3199	CG	LEU	254			125.211	1.00 21.49	В	Ċ
	ATOM	3200		LEU	254			124.364	1.00 20.44	В	C
	ATOM	3201		LEU	254			126.672	1.00 20.62	В	C
20	MOTA	3202		LEU	254						
20			C					125.029	1.00 26.84	В	C
	ATOM	3203	0	LEU	254			124.187	1.00 26.79	В	0
	ATOM	3204	N	GLN	255			125.514	1.00 27.67	В	N
	MOTA	3205	CA	GLN	255			125.037	1.00 29.47	В	С
	MOTA	3206	CB	GLN	255			126.203	1.00 31.78	В	C
25	MOTA	3207	CG	GLN	255			127.263	1.00 36.11	В	С
	MOTA	3208	CD	GLN	255	17.854	-80.644	128.518	1.00 38.94	В	C
	ATOM	3209	OE1	GLN	255	18.321	-80.015	129.476	1.00 40.55	В	0
	ATOM	3210	NE2	GLN	255			128.523	1.00 39.29	В	N
	ATOM	3211	С	GLN	255			123.952	1.00 28.90	В	C
30	MOTA	3212	Ö	GLN	255			123.954	1.00 28.37	В	Ö
00	ATOM	3213	N	GLU	256			123.013	1.00 28.82	В	N
	ATOM	3214	CA	GLU	256						
								121.922	1.00 29.56	В	C
	ATOM	3215	CB	GLU	256			121.102	1.00 30.58	В	C
25	MOTA	3216	CG	GLU	256			120.205	1.00 32.90	В	С
35	MOTA	3217	CD	GLU	256			119.163	1.00 34.07	В	C
	MOTA	3218		GLU	256			119.514	1.00 35.05	В	0
	MOTA	3219	OE2	GLU	256			117.995	1.00 35.39	В	0
	MOTA	3220	С	GLU	256	14.658	-83.154	122.274	1.00 29.06	В	C
	MOTA	3221	0	GLU	256	13.587	-83.160	121.677	1.00 29.72	В	0
40	ATOM	3222	N	PRO	257	14.935	-84.051	123.232	1.00 28.49	В	N
	ATOM	3223	CD	PRO	257			124.019	1.00 28.69	В	C
	ATOM	3224	CA	PRO	257			123.552	1.00 27.78	В	Č
	ATOM	3225	СВ	PRO	257			124.647	1.00 28.06	В	Č
	ATOM	3226	CG	PRO	257			125.201	1.00 29.28		C
45	ATOM	3227	C		257					В	
43				PRO				123.958	1.00 26.26	В	C
	ATOM	3228	0	PRO	257			123.806	1.00 25.48	В	0
	ATOM	3229	N	GLU	258			124.466	1.00 25.25	В	N
	MOTA	3230	CA	GLU	258			124.851	1.00 24.25	В	С
	ATOM	3231	CB	GLU	258			125.831	1.00 24.97	В	C
50	ATOM	3232	CG	GLU	258	12.345	-81.953	127.069	1.00 25.66	В	C
	ATOM	3233	CD	GLU	258	12.787	-80.879	128.029	1.00 25.87	В	С
	ATOM	3234		GLU	258			127.581	1.00 25.45	В	Ō
	ATOM	3235		GLU	. 258			129.240	1.00 25.58	В	Ö
	ATOM	3236	C	GLU	258			123.587	1.00 23.02	В	C
55	ATOM	3237	Ö	GLU	258			123.367	1.00 23.02	В	
	MOTA	3238	N	TYR	259			123.442			0
		3239							1.00 21.21	В	N
	ATOM		CA	TYR	259			121.399	1.00 20.28	В	C
	MOTA	3240	CB	TYR	259	11.939	-80.378	120.527	1.00 18.72	В	С

	ATOM	3241	CG	TYR	259	12.113	-78.882	120.686	1.00 18.66	В	С
	MOTA	3242	CD1		259	11.074	-77.997	120.378	1.00 18.05	В	С
	MOTA	3243			259		-76.616		1.00 17.04	В	С
_	MOTA	3244	CD2		259		-78.346		1.00 16.85	В	С
5	ATOM	3245	CE2		259		-76.976		1.00 15.76	В	С
	MOTA	3246	CZ	TYR	259		-76.114		1.00 17.18	В	C
	ATOM	3247	ОН	TYR	259		-74.750		1.00 16.41	В	0
	ATOM	3248	C	TYR	259		-82.195		1.00 20.33	В	C
10	MOTA MOTA	3249 3250	O N	TYR VAL	259 260		-82.123		1.00 19.27	В	0
10	ATOM	3250	CA	VAL	260		-83.289 -84.498		1.00 20.16	В	N
	ATOM	3252	CB	VAL	260		-85.535		1.00 22.13 1.00 24.17	B B	C
	ATOM	3253	CG1		260		-86.882		1.00 24.17	В	C
	MOTA	3254		VAL	260		-85.088		1.00 25.03	В	C
15	ATOM	3255	C	VAL	260		-85.119		1.00 23.03	В	C
	ATOM	3256	ō	VAL	260		-85.541		1.00 21.90	В	Ö
	ATOM	3257	N	LEU	261		-85.181		1.00 21.23	В	N
	MOTA	3258	CA	LEU	261		-85.746		1.00 22.69	В	C
	MOTA	3259	CB	LEU	261		-85.912		1.00 22.40	В	С
20	ATOM	3260	CG	LEU	261		-87.127		1.00 22.56	В	С
	MOTA	3261	CD1	LEU	261	9.560	-87.121	125.861	1.00 23.42	В	C
	MOTA	3262	CD2	LEU	261	8.419	-88.390	124.019	1.00 21.06	В	C
	MOTA	3263	С	LEU	261		-84.869		1.00 22.97	В	С
	MOTA	3264	0	LEU	261		-85.365		1.00 23.28	В	0
25	ATOM	3265	N	LEU	262		-83.565		1.00 22.71	В	N
	MOTA	3266	CA	LEU	262		-82.611		1.00 23.51	В	C
	MOTA	3267	CB	LEU	262		-81.194		1.00 24.71	В	С
	ATOM	3268	CG	LEU	262		-80.042		1.00 27.53	В	C
20	MOTA	3269		LEU	262		-80.278		1.00 27.55	В	C
30	ATOM	3270		LEU	262		-78.733		1.00 27.91	В	C
	ATOM	3271	C	LEU	262		-82.889		1.00 23.38	В	С
	MOTA	3272	0	LEU	262		-82.918 -83.100		1.00 22.75	В	0
	MOTA MOTA	3273 3274	N CA	ALA ALA	263 263		-83.100 -83.399		1.00 22.06	В	N
35	ATOM	3274	CB	ALA	263		-83.499		1.00 22.18 1.00 20.60	В	C
33	ATOM	3276	C	ALA	263			117.283	1.00 20.80	В	C
	ATOM	3277	0	ALA	263			117.585	1.00 22.24	B B	0
	ATOM	3278	N	ALA	264			118.965	1.00 22.01	В	N
	ATOM	3279	CA	ALA	264			119.073	1.00 22.99	В	C
40	ATOM	3280	СВ	ALA	264			119.905	1.00 23.25	В	C
	ATOM	3281	c	ALA	264			119.707	1.00 23.31	В	Č
	ATOM	3282	0	ALA	264		-87.538		1.00 23.47	В	ō
	ATOM	3283	N	MET	265			120.702	1.00 23.39	В	N
	ATOM	3284	CA	MET	265	2.152	-85.774	121.359	1.00 24.89	В	C
45	ATOM	3285	CB	MET	265	2.326	-84.923	122.617	1.00 26.27	В	C
	ATOM	3286	CG	MET	265	2.960	-85.692	123.761	1.00 28.18	В	C
	MOTA	3287	SD	MET	265	3.160	-84.705	125.239	1.00 30.04	В	S
	MOTA	3288	CE	MET	265			125.842	1.00 28.25	В	C
	MOTA	3289	С	MET	265			120.415	1.00 24.62	В	С
50	MOTA	3290	0	MET	265			120.487	1.00 24.02	В	0
	ATOM	3291	N	ALA	266			119.531	1.00 24.92	В	N
	MOTA	3292	CA	ALA	266			118.550	1.00 25.19	В	C
	ATOM	3293	CB	ALA	266			117.835	1.00 23.17	В	C
55	ATOM	3294	С	ALA	266			117.532	1.00 25.75	В	C
J	ATOM	3295	O	ALA	266			117.156	1.00 25.62	В	0
	ATOM	3296	N	LEU	267			117.092	1.00 25.83	В	N
	ATOM	3297 3298	CA CB	LEU	267 267			116.123	1.00 26.87	В	C
	MOTA	3298	CB	LEU	267	2.296	-01.3//	115.891	1.00 26.43	В	C

	MOTA	3299	CG	LEU	267	2.542	-88.069	114.541	1.00 27.28	В	С
	MOTA	3300	CD1	LEU	267	3.607	-89.134	114.721	1.00 26.00	В	С
	MOTA	3301	CD2	LEU	267		-88.692	113.999	1.00 27.86	В	С
_	MOTA	3302	С	LEU	267		-87.505		1.00 27.40	В	С
5	MOTA	3303	0	LEU	267		-87.741		1.00 27.12	В	0
	MOTA	3304	N	PHE	268		-88.047		1.00 28.25	В	N
	MOTA	3305	CA	PHE	268		-89.006		1.00 30.10	В	С
	ATOM	3306	CB	PHE	268		-90.017		1.00 28.85	В	С
40	MOTA	3307	CG	PHE	268		-90.889	118.483	1.00 28.85	В	С
10	MOTA	3308	CD1		268		-90.857	118.777	1.00 28.83	В	С
	MOTA	3309	CD2		268		-91.709		1.00 28.35	В	C
	ATOM	3310	CE1		268		-91.629		1.00 30.17	В	C
	ATOM ATOM	3311 3312	CE2 CZ		268		-92.486		1.00 29.06	В	C
15	ATOM	3312	C	PHE PHE	268 268		-92.447 -88.427		1.00 28.98	В	С
10	MOTA	3314	0	PHE	268		-88.727		1.00 31.65	В	C
	ATOM	3315	N	SER	269		-87.611		1.00 31.36 1.00 34.17	В	0
	ATOM	3316	CA	SER	269		-87.004		1.00 34.17	B B	N C
	ATOM	3317	СВ	SER	269		-85.581		1.00 36.81	В	C
20	MOTA	3318	OG	SER	269		-84.767		1.00 30.42	В	0
	ATOM	3319	C	SER	269		-87.854		1.00 37.02	В	C
	ATOM	3320	ŏ	SER	269		-87.996		1.00 38.42	В	õ
	ATOM	3321	N	PRO	270		-88.428		1.00 40.51	В	N
	ATOM	3322	CD	PRO	270		-88.310		1.00 40.82	В	C
25	ATOM	3323	CA	PRO	270		-89.275		1.00 41.97	В	Č
	ATOM	3324	CB	PRO	270			120.695	1.00 41.86	В	C
	ATOM	3325	CG	PRO	270	-6.941	-88.839	121.643	1.00 41.56	В	C
	MOTA	3326	С	PRO	270	-8.391	-88.532	118.871	1.00 43.29	В	С
	MOTA	3327	0	PRO	270	-9.311	-89.141	118.325	1.00 43.76	В	0
30	ATOM	3328	N	ASP	271	-8.429	-87.222	119.088	1.00 44.33	В	N
	MOTA	3329	CA	ASP	271	-9.585	-86.424	118.696	1.00 45.68	В	С
	MOTA	3330	CB	ASP	271			119.774	1.00 46.97	В	С
	MOTA	3331	CG	ASP	271			119.928	1.00 48.52	В	С
05	MOTA	3332		ASP	271			119.795	1.00 49.44	В	0
35	MOTA	3333		ASP	271			120.202	1.00 49.05	В	0
	MOTA	3334	С	ASP	271			117.334	1.00 45.97	В	С
	MOTA	3335	0	ASP	271			117.089	1.00 45.91	В	0
	MOTA	3336	N	ARG	272		-86.476		1.00 45.79	В	N
40	ATOM	3337	CA	ARG	272		-85.977		1.00 45.77	В	С
40	ATOM	3338	CB	ARG	272		-86.489		1.00 44.26	В	C
	ATOM	3339	CG	ARG	272			113.998	1.00 41.89	В	C
	ATOM ATOM	3340 3341	CD NE	ARG ARG	272 272			114.976 115.076	1.00 39.65	В	C
	ATOM	3342	CZ	ARG	272			115.076	1.00 38.37 1.00 37.88	В	N
45	ATOM	3343		ARG	272			115.876	1.00 37.88	B B	C
	ATOM	3344		ARG	272			115.565	1.00 37.41	В	N N
	ATOM	3345	C	ARG	272			114.182	1.00 36.36	В	C
	ATOM	3346	ŏ	ARG	272			114.135	1.00 46.53	В	0
	ATOM	3347	N	PRO	273			113.432	1.00 47.81	В	N
50	ATOM	3348	CD	PRO	273			113.255	1.00 48.15	В	C
	ATOM	3349	CA	PRO	273			112.525	1.00 48.97	В	C
	ATOM	3350	CB	PRO	273			111.772	1.00 48.79	В	C
	ATOM	3351	CG	PRO	273			111.838	1.00 48.65	В	c
	ATOM	3352	C	PRO	273			111.594	1.00 49.83	В	Č
55	MOTA	3353	0	PRO	273			110.890	1.00 49.98	В	ō
	MOTA	3354	N	GLY	274			111.615	1.00 50.41	В	N
	MOTA	3355	CA	GLY	274			110.769	1.00 51.30	В	C
	ATOM	3356	С	GLY	274			111.443	1.00 52.28	В	C

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	3.0014	2257	_	OT 12	074	10 267	01 577	110 006	1 00 50 01	_	^
	MOTA	3357	0	GLY	274	-10.367			1.00 52.21	В	0
	MOTA	3358	N	VAL	275	-10.234			1.00 53.24	В	N
	MOTA	3359	CA	VAL	275		-91.500		1.00 54.29	В	С
_	ATOM	3360	CB	VAL	275		-90.984		1.00 54.20	В	С
5	MOTA	3361	CG1		275		-90.054		1.00 54.52	В	С
	ATOM	3362	CG2	VAL	275	-9.742	-90.268	115.621	1.00 54.74	В	С
	MOTA	3363	С	VAL	275	-10.525	-92.604	113.886	1.00 54.76	В	С
	MOTA	3364	0	VAL	275	-11.548	-92.348	114.524	1.00 54.98	В	0
	MOTA	3365	N	THR	276	-10.170	-93.835	113.539	1.00 55.29	В	N
10	ATOM	3366	CA	THR	276		-95.014		1.00 55.88	В	С
	ATOM	3367	СВ	THR	276		-96.152		1.00 55.89	В	Ċ
	MOTA	3368		THR	276		-95.665		1.00 55.88	В	ŏ
	ATOM	3369	CG2	THR	276		-97.315		1.00 56.57	В	Ċ
	ATOM	3370	C	THR	276		-95.527		1.00 56.10	В	C
15	MOTA	3371	0	THR	276		-95.625		1.00 56.37		
13		3372			277		-95.848			В	0
	MOTA		N	GLN					1.00 56.20	В	N
	ATOM	3373	CA	GLN	277		-96.374		1.00 56.61	В	C
	MOTA	3374	CB	GLN	277		-97.051		1.00 57.07	В	С
00	MOTA	3375	CG	GLN	277		-98.557		1.00 58.02	В	С
20	MOTA	3376	CD	GLN	277		-98.975		1.00 58.03	В	С
	MOTA	3377	OE1		277		-98.484		1.00 59.02	В	0
	MOTA	3378	NE2		277		-99.895		1.00 58.06	В	N
	MOTA	3379	С	GLN	277	-8.459	-95.320	117.707	1.00 56.48	В	С
	MOTA	3380	0	GLN	277	-7.325	-95.088	118.130	1.00 56.27	В	0
25	MOTA	3381	N	ARG	278	-9.542	-94.700	118.172	1.00 56.44	В	N
	MOTA	3382	CA	ARG	278	-9.443	-93.672	119.205	1.00 56.32	В	C
	MOTA	3383	СВ	ARG	278	-10.828	-93.134	119.574	1.00 57.69	В	С
	MOTA	3384	CG	ARG	278			118.495	1.00 59.91	В	C
	ATOM	3385	CD	ARG	278			119.105	1.00 62.28	В	Ċ
30	ATOM	3386	NE	ARG	278			119.921	1.00 64.42	В	N
	ATOM	3387	CZ	ARG	278			120.534	1.00 65.26	В	Ĉ
	ATOM	3388		ARG	278			120.436	1.00 65.68	В	N
	MOTA	3389	NH2		278			121.243	1.00 65.69	В	N
	ATOM	3390	C	ARG	278			120.467	1.00 55.50	В	Ç
35	ATOM	3391	0	ARG	278			120.467	1.00 55.30	В	
33				ASP	279			120.830			0
	ATOM	3392	N						1.00 54.66	В	N
	MOTA	3393	CA	ASP	279			122.337	1.00 54.08	В	C
	ATOM	3394	CB	ASP	279			122.869	1.00 55.13	В	C
40	MOTA	3395	CG	ASP	279			123.205	1.00 56.06	В	С
40	MOTA	3396		ASP	279	-11.079		123.898	1.00 56.83	В	0
	MOTA	3397		ASP	279			122.782	1.00 56.30	В	0
	MOTA	3398	С	ASP	279			122.102	1.00 53.02	В	С
	MOTA	3399	0	ASP	279			122.928	1.00 52.50	В	0
	MOTA	3400	N	GLU	280			120.971	1.00 52.25	В	N
45	MOTA	3401	CA	GLU	280	-5.580	-97.151	120.607	1.00 51.45	В	С
	MOTA	3402	CB	GLU	280	-5.605	-97.809	119.221	1.00 53.05	В	С
	MOTA	3403	CG	GLU	280	-6.329	-99.165	119.136	1.00 55.94	В	С
	ATOM	3404	CD	GLU	280	-7.831	-99.090	119.421	1.00 57.54	В	С
	MOTA	3405	OE1	GLU	280	-8.530		118.794	1.00 57.75	В	0
50	MOTA	3406	OE2		280	-8.315		120.271	1.00 59.21	В	0
	ATOM	3407	C	GLU	280			120.605	1.00 49.94	В	Č
	ATOM	3408	Ö	GLU	280			121.211	1.00 49.12	В	0
	ATOM	3409	N	ILE	281			119.919	1.00 48.25	В	
	MOTA	3410	CA	ILE	281	-4.146		119.919			N
55								119.827	1.00 46.84	В	С
33	ATOM	3411	CB	ILE	281				1.00 46.36	В	C
	ATOM	3412		! ILE	281			118.645	1.00 45.98	В	C
	ATOM	3413		ILE	281			117.334	1.00 45.32	В	C
	MOTA	3414	CD1	ILE	281	-4.913	-92.803	116.116	1.00 45.69	В	С

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	MOTA	3415	С	ILE	281		-92.912		1.00 46.37	В	C
	MOTA	3416	0	ILE	281		-92.248		1.00 46.02	В	0
	MOTA	3417	N	ASP	282		-92.986		1.00 46.17	В	Ŋ
_	MOTA	3418	CA	ASP	282		-92.272		1.00 45.94	В	С
5	MOTA	3419	CB	ASP	282		-92.451		1.00 46.56	В	C
	MOTA	3420	CG	ASP	282	-6.918	-91.618	125.038	1.00 48.23	В	С
	MOTA	3421	OD1	ASP	282	-6.723	-90.381		1.00 48.42	В	0
	MOTA	3422	OD2	ASP	282	-7.275	-92.197	126.089	1.00 48.95	В	0
	ATOM	3423	С	ASP	282			124.134	1.00 45.01	В	С
10	MOTA	3424	0	ASP	282			124.864	1.00 45.05	В	0
	MOTA	3425	N	GLN	283	-4.028	-94.088	124.129	1.00 44.59	В	N
	MOTA	3426	CA	GLN	283	-3.018	-94.696	124.994	1.00 43.67	В	С
	ATOM	3427	CB	GLN	283	-3.177	-96.220	124.999	1.00 45.52	В	C
	ATOM	3428	CG	GLN	283	-4.438	-96.701	125.705	1.00 48.81	В	С
15	ATOM	3429	CD	GLN	283	-4.612	-98.206	125.633	1.00 51.03	В	C
	MOTA	3430	OE1	GLN	283	-3.712	-98.967	126.000	1.00 52.12	В	0
	MOTA	3431	NE2	GLN	283	-5.778	-98.646	125.162	1.00 51.60	В	N
	MOTA	3432	С	GLN	283	-1.601	-94.328	124.550	1.00 41.87	В	С
	MOTA	3433	0	GLN	283	-0.710	-94.154	125.376	1.00 40.63	В	0
20	MOTA	3434	N	LEU	284	-1.395	-94.218	123.243	1.00 40.30	В	N
	ATOM	3435	CA	LEU	284			122.718	1.00 39.29	В	C
	MOTA	3436	СВ	LEU	284	-0.092	-93.918	121.185	1.00 39.37	В	C
	ATOM	3437	CG	LEU	284			120.550	1.00 39.93	В	C
	ATOM	3438	CD1	LEU	284			121.249	1.00 40.63	В	C
25	ATOM	3439	CD2	LEU	284			119.067	1.00 38.88	В	C
	ATOM	3440	С	LEU	284			123.188	1.00 37.95	В	C
	MOTA	3441	0	LEU	284			123.638	1.00 37.80	В	ō
	MOTA	3442	N	GLN	285			123.092	1.00 36.96	В	N
	MOTA	3443	CA	GLN	285			123.529	1.00 36.82	В	C
30	MOTA	3444	CB	GLN	285			123.239	1.00 37.92	В	Ċ
	ATOM	3445	CG	GLN	285			123.761	1.00 39.28	В	č
	ATOM	3446	CD	GLN	285			123.334	1.00 40.09	В	Ċ
	ATOM	3447	OE1		285			122.218	1.00 41.73	В	ō
	ATOM	3448	NE2		285			124.211	1.00 40.92	В	N
35	ATOM	3449	С	GLN	285			125.016	1.00 36.21	В	C
	ATOM	3450	ō	GLN	285			125.436	1.00 34.84	В	ŏ
	ATOM	3451	N	GLU	286			125.810	1.00 35.91	В	Ŋ
	ATOM	3452	CA	GLU	286			127.249	1.00 36.06	В	C
	ATOM	3453	СВ	GLU	286			127.937	1.00 38.35	В	č
40	ATOM	3454	CG	GLU	286			129.446	1.00 42.49	В	c
. •	ATOM	3455	CD	GLU	286			130.124	1.00 44.67	В	C
	ATOM	3456		GLU	286			129.837	1.00 46.65		Ö
	ATOM	3457		GLU	286			130.950	1.00 46.57	В	ŏ
	ATOM	3458	C	GLU	286			127.499	1.00 35.04	В	c
45	ATOM	3459	Ö	GLU	286			128.387	1.00 34.20	В	Ö
	ATOM	3460	N	GLU	287			126.708	1.00 34.29	В	N
	ATOM	3461	CA	GLU	287			126.823	1.00 33.78	В	C
	ATOM	3462	CB	GLU	287			125.851	1.00 35.48	В	C
	ATOM	3463	CG	GLU	287			125.864	1.00 38.18	В	C
50	ATOM	3464	CD	GLU	287			124.964	1.00 40.29	В	C
00	ATOM	3465		GLU	287			124.904	1.00 41.09	В	0
	MOTA	3466		GLU	287			. 124.354	1.00 41.61	В	0
	ATOM	3467	C	GLU	287			126.506	1.00 31.01		
	MOTA	3468	o	GLU	287			120.306	1.00 31.95	B B	С О
55	ATOM	3469	N	MET	288			127.137	1.00 31.00	В	
-	ATOM	3470	CA	MET	288			3 125.499 3 125.129	1.00 30.97		N
	ATOM	3471	CB	MET	288			123.129		В	C
	MOTA	3472	CG	MET	288			123.883	1.00 30.43	В	C
	AION	J414	CG	PAE I	200	3.43/	-90.031	122.645	1.00 30.98	В	С

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ATOM 3474 CP MET 288 4.901 -90.867 122.003 1.00 33.03 AND ATOM 3475 CP MET 288 4.251 -88.919 126.300 1.00 22.88 ATOM 3476 CO MET 288 4.251 -88.919 126.501 1.00 28.89 ATOM 3476 CO MET 288 5.310 -88.489 126.751 1.00 28.28 SATOM 3478 CA ALA 289 3.065 -88.568 126.786 1.00 28.24 ATOM 3478 CA ALA 289 3.065 -88.568 126.786 1.00 28.27 ATOM 3478 CA ALA 289 1.441 -87.415 128.195 1.00 28.27 ATOM 3480 CA ALA 289 1.441 -87.415 128.195 1.00 28.77 ATOM 3481 CA ALA 289 3.674 -88.059 129.146 1.00 28.77 ATOM 3481 CA ALA 289 3.674 -88.059 129.146 1.00 28.77 ATOM 3482 N LEU 290 3.555 -89.333 129.511 1.00 29.19 ATOM 3483 CA LEU 290 4.248 -89.850 130.688 1.00 29.79 ATOM 3483 CA LEU 290 3.786 -91.271 131.019 1.00 31.00 ATOM 3485 CG LEU 290 2.029 -92.895 131.750 1.00 31.00 ATOM 3486 CD1 LEU 290 2.029 -92.895 131.750 1.00 33.94 ATOM 3487 CD2 LEU 290 2.029 -92.895 131.750 1.00 33.94 ATOM 3487 CD LEU 290 5.750 -89.850 130.688 1.00 29.75 ATOM 3489 CD LEU 290 5.750 -89.850 130.688 1.00 29.75 ATOM 3489 CA LEU 290 5.750 -89.850 130.475 1.00 29.25 ATOM 3489 CA LEU 290 6.610 -89.852 131.400 1.00 29.25 ATOM 3489 CA THE 291 7.886 -90.695 127.538 1.00 27.69 ATOM 3491 CA THR 291 7.886 -90.695 127.538 1.00 27.69 ATOM 3493 CG THR 291 7.886 -90.695 127.538 1.00 27.69 ATOM 3493 CG THR 291 7.886 -90.695 127.538 1.00 27.69 ATOM 3495 CO THR 291 9.385 -90.688 127.248 1.00 25.40 ATOM 3495 CG LEU 292 7.916 -88.753 129.901 1.00 27.69 ATOM 3498 CG LEU 292 7.929 -86.873 129.691 1.00 27.69 ATOM 3498 CG LEU 292 7.929 -86.873 129.691 1.00 27.69 ATOM 3498 CG LEU 292 7.929 -86.875 129.591 1.00 25.60 ATOM 3499 CG LEU 292 7.929 -86.875 129.591 1.00 25.60 ATOM 3500 CG LEU 292 7.929 -86.875 129.691 1.00 27.73 ATOM 3500 CG LEU 292 7.929 -86.875 129.691 1.00 27.73 ATOM 3500 CG LEU 292 7.929 -86.875 129.691 1.00 27.73 ATOM 3500 CG LEU 292 7.929 -86.875 129.691 1.00 27.73 ATOM 3500 CG LEU 292 7.929 -86.875 129.691 1.00 27.73 ATOM 3500 CG LEU 292 7.929 -86.875 129.691 1.00 27.73 ATOM 3500 CG LEU 292 7.929 -86.875 133.091 1.00 25.61 ATOM 3500 CG LEU 292 7.929 -86.875 133.091												
ATOM 3475 C MET 288 4.251 -88.919 126.300 1.00 28.89 ATOM 3477 N ALA 289 3.065 -88.568 126.786 1.00 28.24 ATOM 3478 CA ALA 289 2.920 -87.622 127.887 1.00 28.24 ATOM 3480 C BALA 289 3.065 -88.568 126.786 1.00 28.77 ATOM 3481 O ALA 289 3.674 -88.059 129.146 1.00 28.77 ATOM 3481 O ALA 289 3.674 -88.059 129.146 1.00 28.77 ATOM 3482 N LEU 290 3.555 -89.333 129.511 1.00 29.19 ATOM 3483 CA LEU 290 3.555 -89.333 129.511 1.00 29.19 ATOM 3485 CG LEU 290 2.029 -92.895 131.750 1.00 33.04 ATOM 3486 CDL LEU 290 2.029 -92.895 131.750 1.00 33.04 ATOM 3486 CDL LEU 290 2.029 -92.895 131.750 1.00 33.94 ATOM 3488 C LEU 290 2.029 -92.895 131.750 1.00 33.94 ATOM 3489 C LEU 290 2.029 -92.895 131.750 1.00 33.94 ATOM 3489 N THR 291 6.510 -89.582 131.400 1.00 29.29 ATOM 3490 N THR 291 6.510 -89.582 131.400 1.00 29.29 ATOM 3491 CA THR 291 7.609 -90.169 128.959 1.00 27.56 ATOM 3493 OG1 THR 291 7.866 -90.695 127.538 1.00 27.59 ATOM 3496 N THR 291 7.866 -90.695 127.538 1.00 27.59 ATOM 3496 N THR 291 7.869 -90.695 127.538 1.00 27.59 ATOM 3497 N LEU 292 7.886 -90.688 127.248 1.00 27.19 ATOM 3498 CA LEU 292 7.886 -90.688 127.248 1.00 27.19 ATOM 3496 N THR 291 7.869 -90.695 127.588 1.00 27.59 ATOM 3497 N LEU 292 7.886 -90.688 127.248 1.00 27.38 ATOM 3496 N LEU 292 7.886 -90.688 127.248 1.00 27.38 ATOM 3496 N LEU 292 7.899 -86.379 128.681 1.00 27.38 ATOM 3497 N LEU 292 7.899 -86.379 128.681 1.00 27.38 ATOM 3498 CA LEU 292 7.899 -86.791 128.856 1.00 27.38 ATOM 3498 CA LEU 292 7.899 -86.791 128.856 1.00 27.38 ATOM 3497 N LEU 292 7.899 -86.791 130.554 1.00 27.38 ATOM 3498 CA LEU 292 7.899 -86.791 130.554 1.00 27.38 ATOM 3498 CA LEU 292 7.899 -86.791 130.554 1.00 27.38 ATOM 3500 CG LEU 292 7.899 -86.791 130.554 1.00 27.37 ATOM 3500 CG LEU 292 7.899 -86.791 130.554 1.00 27.37 ATOM 3500 CG LEU 292 8.890 -83.708 127.599 1.00 27.37 ATOM 3500 C S LEU 292 7.996 -87.708 130.554 1.00 37.79 ATOM 3500 C S LEU 292 7.996 -87.708 130.554 1.00 37.79 ATOM 3500 C S LEU 292 7.996 -87.708 130.0554 1.00 37.75 ATOM 3500 C S LEU 292 8.895 -89.60 130.155 1.00 37.75 ATOM											В	S
5         ATOM         34776         N         MET         288         5.310 - 88.489   126.751         1.00   28.59           ATOM         3478         CA         ALA         289         3.065 - 88.586   126.786   1.00   28.40           ATOM         3480         CA         ALA         289         1.441 - 87.415   128.195   1.00   27.31           10         ATOM         3480         C         ALA         289         4.366 - 87.248   129.777   1.00   28.99           10         ATOM         3481         O         ALA         289         4.356 - 87.248   129.777   1.00   28.99           10         ATOM         3482   CA         LEU         290   4.248 - 89.850   130.688   1.00   29.14           ATOM         3485   CG         LEU         290   2.366 - 91.417   131.574   1.00   31.01           ATOM         3487   CDI   LEU         290   2.259 - 90.677   132.908   1.00   34.21           ATOM         3488   CDI   LEU   290   2.259 - 90.677   132.908   1.00   34.21           ATOM         3487   CDI   TRIP   291   290   2.259   20.677   28.955   130.475   1.00   29.25           ATOM         3495   CDI   TRIP   291   290   2.259   290.677   322.908   1.00   27.56           ATOM         3495   CDI   TRIP   291   290   2.259   290.677   322.908   1.00   27.56           ATOM         3495   CDI   TRIP   291   290   2											В	С
S											В	С
ATOM	_										В	0
ATOM	5										В	N
ATOM											В	C
10 ATOM 3481 O ALA 289											В	С
10											В	С
ATOM	40										В	0
ATOM	10										В	N
ATOM											В	С
ATOM											В	С
15											В	C
ATOM 3488   C   LEU 290   5.750   -89.850   130.475   1.00   29.25											B	С
ATOM 3489 O LEU 290 6.510 -89.582 131.400 1.00 29.29 ATOM 3491 CA THR 291 7.609 -90.169 128.959 1.00 27.56 ATOM 3492 CB THR 291 7.886 -90.695 127.538 1.00 27.59 ATOM 3493 OGI THR 291 7.886 -90.695 127.538 1.00 27.69 ATOM 3494 CG2 THR 291 7.381 -92.034 127.419 1.00 27.19 ATOM 3495 C THR 291 9.385 -90.688 127.248 1.00 25.40 ATOM 3496 C THR 291 9.385 -90.688 127.248 1.00 27.38 ATOM 3496 C THR 291 9.385 -90.688 127.248 1.00 27.39 ATOM 3497 N LEU 292 7.410 -87.772 128.586 1.00 26.70 ATOM 3498 CA LEU 292 7.410 -87.772 128.586 1.00 26.70 ATOM 3499 CB LEU 292 7.829 -86.378 128.681 1.00 27.05 ATOM 3499 CB LEU 292 7.829 -86.378 128.681 1.00 25.96 ATOM 3500 CG LEU 292 7.139 -83.963 127.939 1.00 25.96 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 23.77 ATOM 3503 C LEU 292 6.202 -83.244 126.966 1.00 24.54 ATOM 3503 C LEU 292 7.916 -85.966 130.151 1.00 27.73 ATOM 3505 N GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3506 CA GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3507 CE GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3508 CG GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3508 CG GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 NCEI GLN 293 8.155 -86.547 132.675 1.00 35.44 ATOM 3511 NE2 GLN 293 8.155 -86.547 132.675 1.00 35.44 ATOM 3514 N SER 294 9.550 -90.053 132.962 1.00 32.77 ATOM 3514 N SER 294 9.550 -90.053 132.962 1.00 32.77 ATOM 3514 N SER 294 9.550 -90.053 132.962 1.00 32.77 ATOM 3514 N SER 294 9.550 -90.053 132.962 1.00 32.77 ATOM 3515 CA SER 294 9.550 -90.053 132.962 1.00 32.71 ATOM 3514 N SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3515 CB SER 294 9.550 -90.053 133.900 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 133.900 1.00 34.78 ATOM 3517 CG SER 294 10.808 -87.913 133.090 1.00 34.59 ATOM 3520 N TYR 295 12.175 -86.954 133.351 1.00 34.55 ATOM 3521 CA TYR 295 12.175 -86.954 133.351 1.00 34.55 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3523 CG TYR 295 13.357 -86.155 129.263 1.00 33.57 ATOM 3529 CD TYR 295 15.550 -86.259 128.260 1.00 32.55	15										В	С
ATOM 3490 N THR 291 6.183 -90.158 129.255 1.00 28.88 ATOM 3491 CA THR 291 7.609 -90.169 128.959 1.00 27.569 ATOM 3493 OG1 THR 291 7.886 -90.695 127.538 1.00 27.69 ATOM 3493 OG1 THR 291 7.381 -92.034 127.419 1.00 27.19 ATOM 3494 CG2 THR 291 9.385 -90.688 127.248 1.00 27.38 ATOM 3495 C THR 291 8.159 -88.753 129.090 1.00 27.38 ATOM 3496 CO THR 291 9.243 -88.554 129.637 1.00 26.70 ATOM 3497 N LEU 292 7.410 -87.772 128.586 1.00 26.91 ATOM 3498 CA LEU 292 7.829 -86.378 128.681 1.00 27.05 ATOM 3499 CG LEU 292 7.829 -86.378 128.681 1.00 25.96 ATOM 3499 CG LEU 292 7.829 -86.378 128.681 1.00 25.96 ATOM 3500 CG LEU 292 7.139 -83.963 127.939 1.00 25.61 ATOM 3503 C LEU 292 8.590 -83.708 127.580 1.00 23.77 ATOM 3503 C LEU 292 8.590 -83.708 127.580 1.00 23.77 ATOM 3505 CG LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3505 CG GLEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3505 CG GLN 293 6.917 -86.338 130.949 1.00 26.74 ATOM 3505 CG GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3505 CG GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3505 CG GLN 293 6.917 -86.338 130.949 1.00 33.01 ATOM 3508 CG GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3508 CG GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3511 NE2 GLN 293 3.167 -85.871 132.675 1.00 32.77 ATOM 3513 CC GLN 293 3.167 -86.594 133.059 1.00 33.31 ATOM 3514 NS CR 294 8.356 -87.901 132.881 1.00 37.51 ATOM 3515 CA SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3516 CS SER 294 9.550 -90.053 133.900 1.00 33.57 ATOM 3517 CG SER 294 9.550 -90.053 133.900 1.00 34.88 ATOM 3520 N TYR 295 12.175 -86.954 131.331 1.00 35.00 36.18 ATOM 3520 N TYR 295 12.175 -86.954 131.331 1.00 35.00 36.18 ATOM 3523 CG TYR 295 12.175 -86.694 131.331 1.00 35.00 36.18 ATOM 3520 CD TYR 295 12.175 -86.694 131.331 1.00 35.00 36.18 ATOM 3520 CD TYR 295 12.175 -86.6954 131.331 1.00 35.00 36.18 ATOM 3520 CD TYR 295 12.175 -86.6954 131.331 1.00 35.00 36.18 ATOM 3520 CD TYR 295 12.175 -86.6954 131.331 1.00 35.00 36.18 ATOM 3520 CD TYR 295 12.175 -86.6954 131.331 1.00 33.77 ATOM 3520 CD TYR 295 13.3467 -84.7661 133.3947 1.0										1.00 29.25	В	С
ATOM   3491										1.00 29.29	В	0
ATOM				N	THR					1.00 28.88	В	N
ATOM 3493 OG1 THR 291 7.381 -92.034 127.419 1.00 27.19 ATOM 3495 C THR 291 9.385 -90.688 127.248 1.00 25.48 ATOM 3495 C THR 291 8.159 -88.753 129.090 1.00 27.38 ATOM 3496 O THR 291 9.243 -88.554 129.637 1.00 26.70 ATOM 3497 N LEU 292 7.410 -87.772 128.586 1.00 26.70 ATOM 3498 CA LEU 292 7.829 -86.378 128.6861 1.00 27.05 ATOM 3499 CB LEU 292 7.829 -86.378 128.6861 1.00 27.05 ATOM 3501 CD1 LEU 292 7.139 -83.963 127.939 1.00 25.96 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 23.77 ATOM 3503 C LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3504 O LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3505 N GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3506 CA GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3507 CB GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3508 CG GLN 293 5.663 -86.564 133.059 1.00 35.44 ATOM 3509 CD GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OB1 GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OB1 GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OBL GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OBL GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3515 CA SER 294 8.356 -87.901 132.881 1.00 37.29 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 33.57 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3518 C SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3518 C SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3520 N TYR 295 12.175 -86.954 131.331 1.00 37.51 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3523 CG TYR 295 12.175 -86.954 131.331 1.00 34.20 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 34.20 ATOM 3525 CE1 TYR 295 13.357 -86.115 129.263 1.00 34.78 ATOM 3526 CD2 TYR 295 13.357 -86.854 131.331 1.00 32.77 ATOM 3526 CD2 TYR 295 13.357 -86.854 131.331 1.00 32.77 ATOM 3527 CE2 TYR 295 13.357 -86.854 131.331 1.00 32.75 ATOM 3528 CC2 TYR 295 15.550 -86.6259 128.260 1.00 33.37 ATOM 3529 CH TYR 295 15.550 -86.6259 128.260 1.00 33.57 ATOM 3529 CH TYR 295 15.550 -86.6259 128.260 1.00 33.54		MOTA	3491	CA	THR	291				1.00 27.56	В	С
ATOM 3494 CG2 THR 291 9.385 -90.688 127.248 1.00 25.40 ATOM 3495 C THR 291 8.159 -88.753 129.090 1.00 27.38 ATOM 3496 O THR 291 9.243 -88.554 129.637 1.00 26.70 26.70 ATOM 3498 CA LEU 292 7.410 -87.772 128.586 1.00 26.91 ATOM 3499 CB LEU 292 7.829 -86.378 128.681 1.00 27.05 ATOM 3500 CG LEU 292 7.829 -86.378 128.681 1.00 25.661 ATOM 3501 CD1 LEU 292 8.590 -83.768 127.939 1.00 25.661 ATOM 3502 CD2 LEU 292 6.202 -83.244 126.966 1.00 24.54 ATOM 3503 C LEU 292 6.202 -83.244 126.966 1.00 24.54 ATOM 3503 C LEU 292 8.875 -85.308 130.554 1.00 27.73 ATOM 3504 O LEU 292 8.875 -85.308 130.554 1.00 27.73 ATOM 3505 N GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3506 CA GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3506 CA GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3508 CG GLN 293 4.367 -85.641 133.059 1.00 33.01 ATOM 3508 CG GLN 293 4.367 -85.641 133.059 1.00 33.01 ATOM 3508 CG GLN 293 3.067 -86.541 132.675 1.00 35.44 ATOM 3511 NE2 GLN 293 3.067 -86.541 133.354 1.00 37.29 ATOM 3511 NE2 GLN 293 3.067 -86.594 133.354 1.00 36.31 ATOM 3511 NE2 GLN 293 8.908 -85.875 133.707 1.00 32.11 ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 32.77 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3517 CG SER 294 8.356 -87.901 132.881 1.00 34.78 ATOM 3517 CG SER 294 8.356 -87.901 132.881 1.00 34.78 ATOM 3517 CG SER 294 8.356 -87.901 132.891 1.00 34.78 ATOM 3518 C SER 294 9.502 -88.602 133.460 1.00 34.78 ATOM 3517 CG SER 294 8.355 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 8.356 -87.901 132.811 1.00 33.57 ATOM 3510 CG TYR 295 10.970 -87.613 131.305 1.00 34.78 ATOM 3510 CG TYR 295 10.970 -87.613 131.305 1.00 34.78 ATOM 3520 CG TYR 295 13.357 -86.789 129.814 1.00 34.25 ATOM 3520 CG TYR 295 13.357 -86.868 128.741 1.00 33.57 ATOM 3522 CB TYR 295 13.467 -84.766 128.811 1.00 32.51 ATOM 3522 CB TYR 295 13.467 -84.766 128.811 1.00 32.51 ATOM 3528 CC TYR 295 13.467 -84.766 128.811 1.00 32.51 ATOM 3528 CC TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CC TYR 295 15.550 -86.259 128.260 1.00 33.454 ATOM 3529 OH TYR 295 15.550 -86.2	20	MOTA	3492	СВ	THR	291					В	С
ATOM 3495 C THR 291 8.159 -88.753 129.090 1.00 27.38 ATOM 3496 O THR 291 9.243 -88.554 129.637 1.00 26.70 ATOM 3497 N LEU 292 7.410 -87.772 128.586 1.00 26.91 ATOM 3498 CA LEU 292 7.829 -86.378 128.681 1.00 27.05 ATOM 3499 CB LEU 292 7.829 -86.378 128.681 1.00 25.96 ATOM 3500 CG LEU 292 7.139 -83.963 127.939 1.00 25.96 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 23.77 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 24.54 ATOM 3503 C LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3504 O LEU 292 8.875 -85.960 130.551 1.00 27.73 ATOM 3505 N GLN 293 6.917 -86.338 130.949 1.00 26.74 ATOM 3506 CA GLN 293 6.917 -86.338 130.949 1.00 26.74 ATOM 3508 CG GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3508 CG GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3508 CG GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3501 OE1 GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OE1 GLN 293 3.167 -85.871 132.675 1.00 37.29 ATOM 3510 OE1 GLN 293 3.167 -86.547 134.588 1.00 36.31 ATOM 3511 NC2 GLN 293 8.958 -85.964 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.958 -85.875 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.958 -85.875 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.958 -85.875 133.051 1.00 32.77 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3515 CG SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3515 CG SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3515 CG SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3515 CG SER 294 9.502 -88.602 133.460 1.00 34.89 ATOM 3512 CG TYR 295 12.127 -86.789 129.811 1.00 34.25 ATOM 3520 CB TYR 295 12.127 -86.789 129.811 1.00 33.37 ATOM 3521 CG TYR 295 12.127 -86.789 129.281 1.00 33.37 ATOM 3522 CB TYR 295 12.127 -86.789 129.281 1.00 33.57 ATOM 3522 CB TYR 295 13.357 -86.151 129.263 1.00 33.57 ATOM 3522 CB TYR 295 13.357 -86.151 129.263 1.00 33.57 ATOM 3522 CB TYR 295 13.487 -84.106 128.811 1.00 32.85 ATOM 3525 CE1 TYR 295 13.487 -84.106 128.811 1.00 32.85 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.550 -86.259 128.2		MOTA	3493	OG1	THR	291				1.00 27.19	В	0
25 ATOM 3496 O THR 291 9.243 -88.554 129.637 1.00 26.70 ATOM 3498 CA LEU 292 7.410 -87.772 128.586 1.00 26.91 ATOM 3499 CB LEU 292 6.847 -85.471 127.928 1.00 25.96 ATOM 3500 CG LEU 292 7.139 -83.963 127.939 1.00 25.61 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 24.54 ATOM 3502 CD2 LEU 292 6.202 -83.244 126.966 1.00 24.54 ATOM 3503 C LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3504 O LEU 292 8.875 -85.308 130.554 1.00 27.73 ATOM 3505 N GLN 293 6.917 -86.338 130.949 1.00 26.74 ATOM 3507 CB GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3508 CG GLN 293 5.663 -86.564 133.059 1.00 33.01 ATOM 3508 CG GLN 293 5.663 -86.564 133.059 1.00 37.29 ATOM 3509 CD GLN 293 3.057 -85.871 132.675 1.00 37.29 ATOM 3510 NEZ GLN 293 3.057 -86.491 133.354 1.00 37.29 ATOM 3511 NEZ GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3513 O GLN 293 3.067 -86.564 133.051 1.00 32.77 ATOM 3514 N SER 294 8.355 -86.594 133.051 1.00 32.77 ATOM 3515 CA SER 294 8.355 -86.594 133.051 1.00 32.77 ATOM 3516 CB SER 294 8.355 -86.594 133.051 1.00 32.77 ATOM 3517 NEZ GLN 293 8.908 -85.875 133.707 1.00 32.11 ATOM 3518 C SER 294 8.356 -87.901 132.881 1.00 36.18 ATOM 3517 CA SER 294 8.356 -87.901 132.881 1.00 34.78 ATOM 3518 C SER 294 8.356 -87.901 132.881 1.00 34.78 ATOM 3518 C SER 294 8.356 -87.901 132.881 1.00 34.78 ATOM 3517 CG SER 294 8.356 -87.901 132.881 1.00 34.78 ATOM 3521 CA TYR 295 10.970 -87.613 131.805 1.00 34.78 ATOM 3521 CA TYR 295 10.970 -87.613 131.805 1.00 34.79 ATOM 3522 CB TYR 295 10.970 -87.613 131.805 1.00 34.79 ATOM 3524 CD1 TYR 295 12.175 -86.586 128.261 1.00 33.37 ATOM 3525 CE1 TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3526 CD2 TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3527 CE2 TYR 295 13.460 -86.868 128.741 1.00 33.17  50 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.57 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.57		MOTA	3494	CG2	THR					1.00 25.40	В	C
25 ATOM 3497 N LEU 292 7.410 -87.772 128.586 1.00 26.91 ATOM 3498 CA LEU 292 7.829 -86.378 128.681 1.00 27.05 ATOM 3499 CB LEU 292 7.829 -86.378 128.681 1.00 27.05 ATOM 3500 CG LEU 292 7.139 -83.963 127.939 1.00 25.661 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 23.77 ATOM 3502 CD2 LEU 292 6.202 -83.244 126.966 1.00 24.54 ATOM 3503 C LEU 292 7.916 -88.960 130.151 1.00 27.73 ATOM 3504 O LEU 292 8.875 -85.308 130.554 1.00 26.74 ATOM 3506 CA GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3507 CB GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3507 CB GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3509 CD GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3501 CD1 GLN 293 6.915 -86.491 133.354 1.00 37.29 ATOM 3501 OEI GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3511 NE2 GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3512 C GLN 293 8.908 -85.875 133.707 1.00 32.71 ATOM 3513 O GLN 293 8.908 -85.875 133.707 1.00 32.71 ATOM 3514 N SER 294 9.502 -88.602 133.460 1.00 32.77 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3517 OG SER 294 8.356 -87.901 132.881 1.00 36.18 ATOM 3518 C SER 294 9.502 -88.602 133.460 1.00 34.78 ATOM 3519 O SER 294 8.355 -90.748 133.351 1.00 34.78 ATOM 3520 N TYR 295 10.970 -87.661 133.947 1.00 34.78 ATOM 3521 CA TYR 295 10.970 -87.661 133.947 1.00 34.78 ATOM 3522 CB TYR 295 10.970 -87.661 133.947 1.00 34.78 ATOM 3523 CG TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 33.37 ATOM 3524 CD1 TYR 295 12.127 -86.789 129.814 1.00 33.37 ATOM 3525 CE1 TYR 295 12.127 -86.868 128.741 1.00 33.57 ATOM 3526 CD2 TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3527 CE2 TYR 295 14.627 -84.106 128.811 1.00 32.75 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.54		MOTA	3495	С	THR	291	8.159	-88.753	129.090	1.00 27.38	В	C
ATOM 3498 CA LEU 292 7.829 -86.378 128.681 1.00 27.05 ATOM 3499 CB LEU 292 6.847 -85.471 127.928 1.00 25.96 ATOM 3500 CG LEU 292 7.139 -83.963 127.939 1.00 25.61 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 24.54 ATOM 3503 C LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3503 C LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3504 O LEU 292 8.875 -85.308 130.554 1.00 27.73 ATOM 3505 N GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3506 CA GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3507 CB GLN 293 5.663 -86.564 133.059 1.00 33.01 ATOM 3508 CG GLN 293 4.367 -85.871 132.675 1.00 35.44 ATOM 3509 CD GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3511 NE2 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3511 NE2 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3511 NE2 GLN 293 3.067 -86.547 133.354 1.00 37.29 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3515 CA SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.89 ATOM 3517 OG SER 294 11.654 -87.661 133.947 1.00 34.50 ATOM 3520 N TYR 295 12.175 -86.954 131.331 1.00 35.00 500 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 500 ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3523 CG TYR 295 12.127 -86.789 129.814 1.00 33.37 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 35.00 33.37 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 35.00 33.37 ATOM 3525 CE1 TYR 295 12.127 -86.868 128.741 1.00 32.51 ATOM 3522 CB TYR 295 12.127 -86.868 128.741 1.00 32.51 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.37 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.37 ATOM 3528 CZ TYR		MOTA	3496	0	THR	291	9.243	-88.554	129.637	1.00 26.70	В	0
ATOM 3500 CG LEU 292 7.139 -83.963 127.928 1.00 25.96 ATOM 3501 CD1 LEU 292 7.139 -83.963 127.939 1.00 25.61 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 23.77 ATOM 3502 CD2 LEU 292 6.202 -83.244 126.966 1.00 24.54 ATOM 3503 C LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3505 N GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3506 CA GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3506 CA GLN 293 6.917 -86.338 130.949 1.00 31.80 ATOM 3506 CA GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3508 CG GLN 293 5.663 -86.564 133.059 1.00 33.01 ATOM 3508 CG GLN 293 4.367 -85.871 132.675 1.00 35.44 ATOM 3509 CD GLN 293 3.151 -86.491 133.354 1.00 36.31 ATOM 3510 OE1 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3511 NE2 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3512 C GLN 293 8.908 -85.875 133.707 1.00 32.11 ATOM 3515 CA SER 294 8.356 -87.901 132.881 1.00 32.57 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3519 O SER 294 10.808 -87.913 133.090 1.00 37.51 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.65 ATOM 3521 CA TYR 295 10.970 -87.661 133.947 1.00 34.65 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 34.65 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 35.00 5.00 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 32.57 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 32.55 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 32.55 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 32.55 ATOM 3524 CD1 TYR 295 12.175 -86.954 131.331 1.00 32.55 ATOM 3526 CD2 TYR 295 12.175 -86.954 131.331 1.00 32.55 ATOM 3526 CD2 TYR 295 12.175 -86.954 131.331 1.00 32.55 ATOM 3528 CZ TYR 295 13.357 -86.115 129.263 1.00 32.76 ATOM 3528 CZ TYR 295 15.550 -86.659 128.256 1.00 33.57 ATOM 3528 CZ TYR 295 15.550 -86.659 128.256 1.00 33.54	25	MOTA	3497	N	LEU	292				1.00 26.91	В	N
ATOM 3500 CG LEU 292 7.139 -83.963 127.939 1.00 25.61 ATOM 3501 CD1 LEU 292 8.590 -83.708 127.580 1.00 23.77 ATOM 3502 CD2 LEU 292 6.202 -83.244 126.966 1.00 24.54 ATOM 3503 C LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3505 N GLN 293 6.917 -86.338 130.554 1.00 26.74 ATOM 3506 CA GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3507 CB GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3508 CG GLN 293 5.663 -86.564 133.059 1.00 33.01 ATOM 3509 CD GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OE1 GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3511 NE2 GLN 293 3.067 -86.547 134.588 1.00 36.29 ATOM 3511 NE2 GLN 293 3.067 -86.547 132.546 1.00 38.33 ATOM 3514 N SER 294 8.355 -86.594 133.051 1.00 32.77 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 33.57 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.78 ATOM 3518 C SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3519 O SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3520 N TYR 295 10.970 -87.661 133.3947 1.00 34.29 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3523 CG TYR 295 12.175 -86.688 128.741 1.00 34.28 ATOM 3524 CD1 TYR 295 13.357 -86.115 129.263 1.00 33.57 ATOM 3526 CD2 TYR 295 13.467 -84.106 128.811 1.00 32.77 ATOM 3526 CD TYR 295 13.467 -84.106 128.811 1.00 32.77 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.255 1.00 33.57 ATOM 3528 CZ TYR 295 15.550 -86.259 128.255 1.00 33.54		MOTA	3498	CA	LEU	292				1.00 27.05	В	C
ATOM   3501   CD1   LEU   292   8.590   -83.708   127.580   1.00   23.77		MOTA	3499	СВ	LEU	292 <sup>°</sup>	6.847	-85.471	127.928	1.00 25.96	В	С
30       ATOM       3502       CD2       LEU       292       6.202       -83.244       126.966       1.00       24.54         ATOM       3503       C       LEU       292       7.916       -85.960       130.151       1.00       27.73         ATOM       3504       O       LEU       292       8.875       -85.308       130.554       1.00       26.74         ATOM       3505       N       GLN       293       6.915       -86.008       132.374       1.00       29.30         ATOM       3506       CA       GLN       293       6.915       -86.008       132.374       1.00       31.80         35       ATOM       3508       CG       GLN       293       4.367       -85.871       132.675       1.00       33.01         ATOM       3510       OEI       GLN       293       3.151       -86.491       133.354       1.00       36.31         ATOM       3511       NEZ       GLN       293       3.151       -86.491       133.354       1.00       36.31         40       ATOM       3512       C       GLN       293       8.155       -86.594       133.051       1.00		MOTA	3500	CG	LEU	292	7.139	-83.963	127.939	1.00 25.61	В	С
ATOM 3503 C LEU 292 7.916 -85.960 130.151 1.00 27.73 ATOM 3504 O LEU 292 8.875 -85.308 130.554 1.00 26.74 ATOM 3505 N GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3506 CA GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3507 CB GLN 293 5.663 -86.564 133.059 1.00 33.01 ATOM 3508 CG GLN 293 4.367 -85.871 132.675 1.00 35.44 ATOM 3509 CD GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OE1 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3511 NE2 GLN 293 2.201 -86.964 133.051 1.00 38.33 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3516 CB SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3518 C SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3518 C SER 294 9.502 -90.053 132.962 1.00 36.18 ATOM 3518 C SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 8.353 -90.748 133.261 1.00 34.78 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3523 CG TYR 295 12.175 -86.954 131.331 1.00 33.37 ATOM 3523 CG TYR 295 12.175 -86.954 131.331 1.00 32.51 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3525 CE1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 13.487 -84.726 129.289 1.00 32.70 ATOM 3526 CD2 TYR 295 13.487 -84.726 129.289 1.00 32.70 ATOM 3528 CC2 TYR 295 15.550 -86.259 128.260 1.00 33.75 ATOM 3528 CC2 TYR 295 15.550 -86.259 128.260 1.00 33.75 ATOM 3528 CC2 TYR 295 15.550 -86.259 128.260 1.00 33.75 ATOM 3528 CC2 TYR 295 15.550 -86.259 128.260 1.00 33.75 ATOM 3528 CC2 TYR 295 15.550 -86.259 128.260 1.00 33.75 ATOM 3528 CC2 TYR 295 15.550 -86.259 128.260 1.00 33.75 ATOM 3529 OH TYR 295 15.550 -86.259 128.250 1.00 33.54 ATOM 3529 OH TYR 295 15.550 -86.259 128.250 1.00 33.54 ATOM 3529 OH TYR 295 15.550 -86.259 128.250 1.00 33.54 ATOM 3529 OH TYR 295 15.550 -86.259 128.250 1.00 33.54		MOTA	3501	CD1	LEU	292	8.590	-83.708	127.580	1.00 23.77	В	С
ATOM 3504 O LEU 292 8.875 -85.308 130.554 1.00 26.74 ATOM 3505 N GLN 293 6.917 -86.338 130.949 1.00 29.30 ATOM 3506 CA GLN 293 6.915 -86.008 132.374 1.00 31.80 ATOM 3507 CB GLN 293 5.663 -86.564 133.059 1.00 33.01 ATOM 3508 CG GLN 293 4.367 -85.871 132.675 1.00 35.44 ATOM 3509 CD GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OE1 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3511 NE2 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.155 -86.594 133.707 1.00 32.77 ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3518 C SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3518 C SER 294 11.654 -87.661 133.947 1.00 34.78 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3524 CD1 TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3525 CE1 TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3526 CD2 TYR 295 13.487 -84.726 129.289 1.00 32.70 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 33.54 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.554 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.554	30	MOTA	3502	CD2	LEU	292	6.202	-83.244	126.966	1.00 24.54	В	C
ATOM 3506 CA GLN 293 6.917 -86.338 130.949 1.00 29.30  ATOM 3506 CA GLN 293 6.915 -86.008 132.374 1.00 31.80  ATOM 3507 CB GLN 293 5.663 -86.564 133.059 1.00 33.01  ATOM 3508 CG GLN 293 4.367 -85.871 132.675 1.00 35.44  ATOM 3509 CD GLN 293 3.151 -86.491 133.354 1.00 36.31  ATOM 3510 OE1 GLN 293 3.067 -86.547 134.588 1.00 36.31  ATOM 3511 NE2 GLN 293 2.201 -86.964 132.546 1.00 38.33  40 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77  ATOM 3513 O GLN 293 8.908 -85.875 133.707 1.00 32.77  ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57  ATOM 3515 CA SER 294 9.550 -90.053 132.962 1.00 36.18  45 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18  ATOM 3518 C SER 294 8.353 -90.748 133.261 1.00 37.51  ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.78  ATOM 3520 N TYR 295 10.970 -87.661 133.947 1.00 34.65  ATOM 3521 CA TYR 295 10.970 -87.661 133.947 1.00 34.65  ATOM 3522 CB TYR 295 12.175 -86.789 129.814 1.00 33.37  ATOM 3523 CG TYR 295 12.127 -86.789 129.814 1.00 33.37  ATOM 3524 CD1 TYR 295 13.357 -86.115 129.263 1.00 33.37  ATOM 3525 CE1 TYR 295 13.487 -84.726 129.289 1.00 32.85  ATOM 3526 CD2 TYR 295 13.487 -84.726 129.289 1.00 32.51  ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.57  ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54  ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54		MOTA	3503	С	LEU	292	7.916	-85.960	130.151		В	С
ATOM   3506   CA   GLN   293   6.915 -86.008   132.374   1.00   31.80		ATOM	3504	0	LEU	292				1.00 26.74	В	0
35         ATOM         3507         CB         GLN         293         5.663         -86.564         133.059         1.00         33.01           ATOM         3508         CG         GLN         293         4.367         -85.871         132.675         1.00         35.44           ATOM         3509         CD         GLN         293         3.151         -86.491         133.354         1.00         37.29           ATOM         3510         OE1         GLN         293         3.067         -86.547         134.588         1.00         36.31           ATOM         3511         NE2         GLN         293         8.155         -86.594         132.546         1.00         38.33           40         ATOM         3512         C         GLN         293         8.155         -86.594         132.546         1.00         38.33           40         ATOM         3512         O         GLN         293         8.155         -86.594         133.051         1.00         32.77           ATOM         3514         N         SER         294         8.356         -87.901         132.881         1.00         33.57           ATOM <t< th=""><th></th><th>MOTA</th><th>3505</th><th>N</th><th>GLN</th><th>293</th><th>6.917</th><th>-86.338</th><th>130.949</th><th>1.00 29.30</th><th>В</th><th>N</th></t<>		MOTA	3505	N	GLN	293	6.917	-86.338	130.949	1.00 29.30	В	N
ATOM 3508 CG GLN 293 4.367 -85.871 132.675 1.00 35.44 ATOM 3509 CD GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OE1 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3511 NE2 GLN 293 2.201 -86.964 132.546 1.00 38.33  40 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.908 -85.875 133.707 1.00 32.11 ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18  45 ATOM 3518 C SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3519 O SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00  50 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 35.00  50 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3525 CE1 TYR 295 13.487 -84.726 129.289 1.00 32.851 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 33.17  55 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54		MOTA	3506	CA	GLN	293				1.00 31.80	В	C
ATOM 3510 OE1 GLN 293 3.151 -86.491 133.354 1.00 37.29 ATOM 3510 OE1 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3511 NE2 GLN 293 2.201 -86.964 132.546 1.00 38.33  40 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.908 -85.875 133.707 1.00 32.71 ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3519 O SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3520 N TYR 295 10.970 -87.661 133.947 1.00 34.65 ATOM 3521 CA TYR 295 10.970 -87.661 133.947 1.00 34.65 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 35.00  50 ATOM 3523 CG TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 15.550 -86.259 128.260 1.00 33.77 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54	35	ATOM	3507	СВ	GLN	293	5.663	-86.564	133.059	1.00 33.01	В	C
ATOM 3510 OE1 GLN 293 3.067 -86.547 134.588 1.00 36.31 ATOM 3511 NE2 GLN 293 2.201 -86.964 132.546 1.00 38.33 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.908 -85.875 133.707 1.00 32.11 ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 50 ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 33.37 ATOM 3525 CE1 TYR 295 13.487 -84.726 129.289 1.00 33.17 55 ATOM 3526 CD2 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.54 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54		ATOM	3508	CG	GLN	293	4.367	-85.871	132.675	1.00 35.44	В	C
40 ATOM 3511 NE2 GLN 293 2.201 -86.964 132.546 1.00 38.33 8.155 -86.594 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.908 -85.875 133.707 1.00 32.11 ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3518 C SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3521 CA TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 35.00 50 ATOM 3522 CB TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3525 CE1 TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3526 CD2 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3527 CE2 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54		ATOM	3509	CD	GLN	293	3.151	-86.491	133.354	1.00 37.29	В	C
40 ATOM 3512 C GLN 293 8.155 -86.594 133.051 1.00 32.77 ATOM 3513 O GLN 293 8.908 -85.875 133.707 1.00 32.11 ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3516 CB SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3523 CG TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3524 CD1 TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3525 CE1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3526 CD2 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 33.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54		MOTA	3510	OE1	GLN	293				1.00 36.31	В	0
ATOM 3513 O GLN 293 8.908 -85.875 133.707 1.00 32.11 ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CEI TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 33.54 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54		MOTA	3511	NE2	GLN	293	2.201	-86.964	132.546	1.00 38.33	В	N
ATOM 3514 N SER 294 8.356 -87.901 132.881 1.00 33.57 ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3523 CG TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3524 CD1 TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3525 CE1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3526 CD2 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3527 CE2 TYR 295 14.406 -86.868 128.741 1.00 33.17  55 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54	40	MOTA	3512	C	GLN	293	8.155	-86.594	133.051	1.00 32.77	В	C
ATOM 3515 CA SER 294 9.502 -88.602 133.460 1.00 34.88 ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.175 -86.789 129.814 1.00 34.20 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3526 CD2 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 33.77 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		ATOM	3513	0	GLN	293	8.908	-85.875	133.707	1.00 32.11	В	0
ATOM 3516 CB SER 294 9.550 -90.053 132.962 1.00 36.18 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54		MOTA		N	SER	294	8.356	-87.901	132.881		В	N
45 ATOM 3517 OG SER 294 8.353 -90.748 133.261 1.00 37.51 ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3526 CD2 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3527 CE2 TYR 295 14.406 -86.868 128.741 1.00 33.17 55 ATOM 3528 CZ TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3529 OH TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		MOTA		CA		294				1.00 34.88	В	C
ATOM 3518 C SER 294 10.808 -87.913 133.090 1.00 34.78 ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17  55 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		MOTA	3516	CB	SER					1.00 36.18	В	C
ATOM 3519 O SER 294 11.654 -87.661 133.947 1.00 34.65 ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00  ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17  55 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15	45	MOTA	3517	OG	SER	294	8.353	-90.748	133.261	1.00 37.51	В	0
ATOM 3520 N TYR 295 10.970 -87.613 131.805 1.00 34.99 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17  55 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		MOTA	3518	С	SER	294	10.808	-87.913	133.090	1.00 34.78	В	С
50 ATOM 3521 CA TYR 295 12.175 -86.954 131.331 1.00 35.00 ATOM 3522 CB TYR 295 12.127 -86.789 129.814 1.00 34.20 ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		ATOM	3519	0	SER	294				1.00 34.65	В	0
50         ATOM         3522         CB         TYR         295         12.127         -86.789         129.814         1.00         34.20           ATOM         3523         CG         TYR         295         13.357         -86.115         129.263         1.00         33.37           ATOM         3524         CD1         TYR         295         13.487         -84.726         129.289         1.00         32.51           ATOM         3525         CE1         TYR         295         14.627         -84.106         128.811         1.00         32.85           ATOM         3526         CD2         TYR         295         14.406         -86.868         128.741         1.00         33.17           55         ATOM         3527         CE2         TYR         295         15.550         -86.259         128.260         1.00         32.70           ATOM         3528         CZ         TYR         295         15.654         -84.879         128.295         1.00         33.54           ATOM         3529         OH         TYR         295         16.779         -84.275         127.784         1.00         34.15		MOTA	3520	N	TYR	295	10.970	-87.613	131.805	1.00 34.99	В	N
ATOM 3523 CG TYR 295 13.357 -86.115 129.263 1.00 33.37 ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		MOTA	3521	CA	TYR	295	12.175	-86.954	131.331	1.00 35.00	В	С
ATOM 3524 CD1 TYR 295 13.487 -84.726 129.289 1.00 32.51 ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15	50	MOTA	3522	CB	TYR	295	12.127	-86.789	129.814	1.00 34.20	В	С
ATOM 3525 CE1 TYR 295 14.627 -84.106 128.811 1.00 32.85 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		MOTA	3523	CG	TYR					1.00 33.37	В	С
55 ATOM 3526 CD2 TYR 295 14.406 -86.868 128.741 1.00 33.17 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		ATOM	3524							1.00 32.51	В	C
55 ATOM 3527 CE2 TYR 295 15.550 -86.259 128.260 1.00 32.70 ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		MOTA	3525	CE1						1.00 32.85	В	С
ATOM 3528 CZ TYR 295 15.654 -84.879 128.295 1.00 33.54 ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15		MOTA	3526	CD2	YYR					1.00 33.17	В	С
ATOM 3529 OH TYR 295 16.779 -84.275 127.784 1.00 34.15	55	MOTA	3527	CE2	YYR					1.00 32.70	В	С
		MOTA		CZ	TYR		15.654	-84.879	128.295	1.00 33.54	В	С
		MOTA		ОН							В	0
		MOTA	3530	C	TYR	295	12.391	-85.598	3 131.993	1.00 35.91	В	

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	ATOM	3531	0	TYR	295	13.510	-85.268	132.379	1.00 35.15	В	0
	MOTA	3532	N	ILE	296		-84.809		1.00 37.71	В	N
	ATOM	3533	CA	ILE	296		-83.492		1.00 40.23	В	C
	MOTA	3534	СВ	ILE	296		-82.705		1.00 39.25	В	C
5	ATOM	3535	CG2		296		-81.424		1.00 39.23		C
	ATOM	3536	CG1		296		-82.378		1.00 38.18	В	
	ATOM	3537	CD1		296		-81.651			В	C
	ATOM	3538	CDI						1.00 38.48	В	C
		3539	0	ILE	296		-83.639		1.00 42.98	В	C
10	MOTA			ILE	296		-82.934		1.00 42.72	В	0
10	ATOM	3540	N	LYS	297		-84.552		1.00 46.09	В	N
	MOTA	3541	CA	LYS	297		-84.799		1.00 49.98	В	C
	ATOM	3542	CB	LYS	297		-85.997		1.00 49.95	В	C
	MOTA	3543	CG	LYS	297		-85.718		1.00 50.90	В	C
15	MOTA	3544	CD	LYS	297		-86.957		1.00 51.71	В	C
13	ATOM	3545	CE	LYS	297		-86.713		1.00 51.90	В	С
	MOTA	3546	NZ	LYS	297		-87.944		1.00 52.46	В	N
	MOTA	3547	C	LYS	297		-85.055		1.00 52.54	В	С
	ATOM	3548	0	LYS	297			137.500	1.00 53.13	В	0
20	ATOM	3549	N	GLY	298			135.919	1.00 55.39	В	N
20	ATOM	3550	CA	GLY	298			136.167	1.00 59.24	В	С
	ATOM	3551	C	GLY	298			135.485	1.00 62.03	В	С
	ATOM	3552	0	GLY	298			135.824	1.00 62.62	В	0
	ATOM	3553	N	GLN	299			134.536	1.00 64.81	В	N
25	ATOM	3554	CA	GLN	299			133.816	1.00 67.60	В	C
25	MOTA	3555	CB	GLN	299			132.688	1.00 67.76	В	С
	MOTA	3556	CG	GLN	299			131.729	1.00 68.45	В	С
	ATOM	3557	CD	GLN	299			131.453	1.00 68.63	В	С
	ATOM	3558		GLN	299			131.236	1.00 68.95	В	0
20	MOTA	3559	NE2		299			131.457	1.00 68.63	В	N
30	ATOM	3560	C	GLN	299			134.697	1.00 69.34	В	C
	MOTA	3561	0	GLN	299			135.518	1.00 69.59	В	0
	MOTA	3562	N	GLN	300			134.495	1.00 71.45	В	N
	ATOM	3563	CA	GLN	300			135.221	1.00 73.26	В	С
25	MOTA	3564	СВ	GLN	300			134.676	1.00 73.69	В	С
35	ATOM	3565	CG	GLN	300			135.213	1.00 74.64	В	С
	ATOM	3566	CD	GLN	300 -	20.675	-84.451	134.846	1.00 75.14	В	C
	ATOM	3567		GLN	300			133.667	1.00 75.32	В	0
	ATOM	3568	NE2		300			135.858	1.00 75.53	В	N
40	ATOM	3569	C	GLN	300			135.204	1.00 74.21	В	C
40	MOTA	3570	0	GLN	300			135.303	1.00 74.60	В	0
	MOTA	3571	N	ARG	301			135.073	1.00 75.01	В	N
	ATOM	3572	CA	ARG	301			135.086	1.00 75.30	В	С
	MOTA	3573	CB	ARG	301			133.858	1.00 75.26	В	C
AE	MOTA	3574	CG	ARG	301			133.901	1.00 75.18	В	С
45	ATOM	3575	CD	ARG	301			132.580	1.00 75.06	В	С
	ATOM	3576	NE	ARG	301			132.349	1.00 74.97	В	N
	MOTA	3577	CZ	ARG	301			131.281	1.00 74.61	В	С
	MOTA	3578		ARG	301			130.317	1.00 74.17	В	N
50	MOTA	3579		ARG	301			131.174	1.00 74.08	В	N
50	ATOM	3580	С	ARG	301			136.384	1.00 75.55	В	C
	MOTA	3581	0	ARG	301			136.615	1.00 75.54	В	0
	ATOM	3582	N	ARG	302			137.240	1.00 75.70	В	N
	MOTA	3583	CA	ARG	302			138.507	1.00 75.72	В	С
EE	ATOM	3584	CB	ARG	302			139.602	1.00 76.38	В	С
55	MOTA	3585	CG	ARG	302			141.018	1.00 77.18	В	С
	ATOM	3586	CD	ARG	302			141.988	1.00 77.70	В	С
	ATOM	3587	NE	ARG	302			142.207	1.00 78.27	В	N
	MOTA	3588	CZ	ARG	302	21.562	-75.290	142.868	1.00 78.26	В	С

ATOM 3590 NH2 ARG 302 22.767 -74.760 143.016 1.00 78.37 B ATOM 3591 C ARG 302 17.762 -75.317 138.574 1.00 75.23 B ATOM 3593 N PRO 303 17.299 -74.803 139.663 1.00 74.61 B ATOM 3593 N PRO 303 17.292 -74.803 139.663 1.00 74.61 B ATOM 3595 CA PRO 303 17.298 -75.150 135.986 1.00 74.54 B ATOM 3595 CA PRO 303 16.273 -73.750 137.667 1.00 73.72 B ATOM 3596 CB PRO 303 16.273 -73.750 137.667 1.00 73.72 B ATOM 3596 CB PRO 303 16.012 -73.215 136.261 1.00 73.72 B ATOM 3596 CB PRO 303 16.012 -73.215 136.261 1.00 72.47 B ATOM 3598 C PRO 303 16.012 -73.579 138.807 1.00 72.47 B ATOM 3599 C PRO 303 15.010 -74.319 138.321 1.00 72.47 B ATOM 3600 N ARG 304 14.920 -75.648 138.330 1.00 71.25 B ATOM 3600 CB ARG 304 14.920 -75.648 138.330 1.00 71.25 B ATOM 3602 CB ARG 304 13.947 -76.484 140.423 1.00 71.29 B ATOM 3603 CG ARG 304 14.921 -77.652 140.877 1.00 72.92 B ATOM 3603 CG ARG 304 14.921 -77.652 140.877 1.00 72.92 B ATOM 3605 CD ARG 304 14.921 -77.652 140.877 1.00 72.92 B ATOM 3605 CC ARG 304 14.821 -77.652 140.877 1.00 75.67 B ATOM 3606 CZ ARG 304 15.103 -79.684 144.027 1.00 76.56 B ATOM 3606 CZ ARG 304 15.103 -79.684 144.027 1.00 76.56 B ATOM 3608 NH2 ARG 304 15.103 -79.684 144.027 1.00 76.34 B ATOM 3608 NH2 ARG 304 15.103 -79.684 144.027 1.00 76.35 B ATOM 3608 NH2 ARG 304 15.103 -79.684 144.027 1.00 76.35 B ATOM 3610 CA ARG 304 15.103 -79.684 144.027 1.00 76.35 B ATOM 3610 CA ARG 304 15.103 -79.684 144.027 1.00 76.35 B ATOM 3610 CA ARG 304 15.103 -79.684 144.027 1.00 76.35 B ATOM 3610 CA ARG 304 15.103 -79.684 144.027 1.00 76.35 B ATOM 3610 CA ARG 304 15.103 -79.684 144.027 1.00 76.35 B ATOM 3610 CA ARG 304 15.103 -79.684 144.027 1.00 76.35 B ATOM 3610 CA ARG 304 17.889 -74.221 135.082 1.00 67.71 B ATOM 3610 CA ARG 304 17.889 -74.221 135.082 1.00 62.41 B ATOM 3610 CA ARG 304 17.889 -74.221 135.082 1.00 62.41 B ATOM 3610 CA ARG 306 R.888 -76.533 138.016 1.00 58.07 B ATOM 3612 CA ARG 306 R.888 -76.543 138.801 1.00 62.41 B ATOM 3612 CA ARG 306 R.888 -76.543 138.801 1.00 62.41 B ATOM 3620 CA ARG 306 R.888 -76.543 138.801 1.00 62.41 B								
ATOM 3599 C ARG 302 17.762 -75.317 138.574 1.00 75.23 B ATOM 3599 N PRO 303 17.202 -74.860 137.430 1.00 75.35 B ATOM 3595 CD PRO 303 17.202 -74.860 137.430 1.00 73.58 B ATOM 3596 CB PRO 303 16.012 -73.215 136.261 1.00 73.58 B ATOM 3596 CB PRO 303 16.012 -73.215 136.261 1.00 73.72 B ATOM 3597 CG PRO 303 16.012 -73.215 136.261 1.00 73.72 B ATOM 3598 C PRO 303 16.012 -73.215 136.261 1.00 73.72 B ATOM 3598 C PRO 303 15.010 -74.319 138.321 1.00 72.47 B ATOM 3598 C PRO 303 15.010 -74.319 138.321 1.00 72.47 B ATOM 3600 N ARG 304 14.920 -75.648 138.330 1.00 72.47 B ATOM 3601 CA ARG 304 13.946 -75.799 138.807 1.00 72.47 B ATOM 3602 CB ARG 304 14.920 -75.648 138.330 1.00 71.25 B ATOM 3603 CG ARG 304 14.673 -77.891 142.376 1.00 72.27 B ATOM 3605 CB ARG 304 14.673 -77.891 142.376 1.00 72.29 B ATOM 3606 CZ ARG 304 14.673 -77.891 142.376 1.00 72.29 B ATOM 3608 NH2 ARG 304 15.013 -79.684 144.907 1.00 72.25 B ATOM 3608 NH2 ARG 304 15.013 -79.684 144.907 1.00 76.25 B ATOM 3608 NH2 ARG 304 15.613 -80.876 144.309 1.00 76.25 B ATOM 3608 NH2 ARG 304 15.613 -80.876 144.300 1.00 66.25 B ATOM 3601 C ARG 304 11.882 -74.999 139.379 1.00 68.11 B ATOM 3611 N ASP 305 11.989 -75.956 137.344 1.00 76.25 B ATOM 3612 CA ASP 305 11.989 -75.956 137.344 1.00 76.25 B ATOM 3613 CB ASP 305 11.989 -75.956 137.344 1.00 64.75 B ATOM 3610 C ARG 304 11.882 -74.999 139.379 1.00 68.11 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 62.77 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 62.77 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 62.77 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 62.77 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3610 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3630 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B		MOTA	3589	NH1	ARG	302		B N
**ATOM** 3599** O.** ARG. **302** 17.499** 74.803** 139.663** 1.00** 74.54** B** ATOM** 3595** CA** PRO** 303** 17.298** 75.150** 135.986** 1.00** 74.54** B** ATOM** 3595** CA** PRO** 303** 16.273** 73.750** 135.986** 1.00** 74.54** B** ATOM** 3595** CA** PRO** 303** 16.273** 73.750** 137.667** 1.00** 73.72** B** ATOM** 3596** CB** PRO** 303** 16.012** 73.215** 136.261** 1.00** 73.72** B** ATOM** 3599** C** PRO** 303** 16.073** 74.451** 135.427** 1.00** 72.47** B** ATOM** 3599** C** PRO** 303** 16.073** 74.451** 135.427** 1.00** 72.47** B** ATOM** 3599** C** PRO** 303** 14.156** 73.579** 138.807** 1.00** 72.47** B** ATOM** 3600** N** ARG. ** 304** 13.796** 76.366** 138.901** 1.00** 72.47** B** ATOM** 3601** CA** ARG. ** 304** 13.796** 76.366** 138.901** 1.00** 71.25** B** ATOM** 3602** CB** ARG. ** 304** 13.796** 76.366** 138.901** 1.00** 71.29** B** ATOM** 3603** CG** ARG. ** 304** 14.821** 777.652** 140.877** 1.00** 72.92** B** ATOM** 3604** CD** ARG. ** 304** 15.207** 7-91.88** 142.794** 1.00** 75.67** B** ATOM** 3606** CZ** ARG. ** 304** 15.207** 7-91.88** 142.794** 1.00** 75.65** B** ATOM** 3608** NH2** ARG. ** 304** 15.503** 7-95.684** 144.027** 1.00** 76.56** B** ATOM** 3608** NH2** ARG. ** 304** 15.613** 808.876** 144.309** 1.00** 76.56** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.562** 1.00** 67.71** B** ATOM** 3610** O** ARG. ** 304** 12.644** 7-5.716** 138.		MOTA			ARG	302		
S		MOTA		С	ARG	302		в С
ATOM	_	MOTA	3592	0	ARG	302	17.499 -74.803 139.663 1.00 75.35	в О
ATOM 3595 CA PRO 303 16.273 -73.750 137.667 1.00 73.58 B ATOM 3596 CB PRO 303 16.012 -73.215 136.261 1.00 74.08 B ATOM 3597 CG PRO 303 16.012 -73.215 136.261 1.00 74.08 B ATOM 3598 C PRO 303 15.010 -74.319 138.321 1.00 72.47 B ATOM 3599 O PRO 303 14.156 -73.579 138.807 1.00 72.47 B ATOM 3600 N ARG 304 14.920 -75.648 138.330 1.00 71.25 B ATOM 3601 CG ARG 304 13.947 -76.484 140.423 1.00 74.08 B ATOM 3602 CB ARG 304 13.947 -76.484 140.423 1.00 71.29 B ATOM 3603 CG ARG 304 13.947 -76.521 140.877 1.00 72.92 B ATOM 3603 CG ARG 304 14.673 -77.881 142.376 1.00 74.32 B ATOM 3603 CC ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3605 NE ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3606 NE ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3606 NE ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3606 NE ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3608 NE ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3608 NE ARG 304 15.207 -79.188 142.794 1.00 76.556 B ATOM 3609 C ARG 304 12.464 -78.995 144.980 1.00 76.556 B ATOM 3609 C ARG 304 12.464 -78.995 144.980 1.00 76.548 B ATOM 3601 N ASP 305 11.989 -75.956 137.344 1.00 64.75 B ATOM 3611 N ASP 305 10.788 -75.16 138.562 1.00 67.71 B ATOM 3612 CB ASP 305 10.789 -74.800 135.594 1.00 61.19 B ATOM 3613 CB ASP 305 10.789 -74.800 135.594 1.00 62.41 B ATOM 3613 CB ASP 305 10.789 -74.800 135.594 1.00 62.41 B ATOM 3613 CB ASP 305 10.789 -74.800 135.594 1.00 62.77 B ATOM 3612 CB ASP 305 9.202 -74.194 133.860 1.00 62.71 B ATOM 3613 CB ASP 305 9.625 -77.382 136.059 1.00 62.77 B ATOM 3612 CB ASP 305 9.625 -77.382 136.059 1.00 62.77 B ATOM 3612 CB ARG 306 7.440 -77.548 138.187 1.00 55.34 B ATOM 3613 CB ASP 305 9.202 -74.194 133.860 1.00 62.71 B ATOM 3612 CB ARG 306 7.440 -77.548 138.187 1.00 55.34 B ATOM 3613 CB ASP 305 9.202 -74.800 135.594 1.00 62.77 B ATOM 3612 CB ARG 306 7.440 -77.548 138.187 1.00 55.34 B ATOM 3620 CB ARG 306 7.440 -77.548 138.187 1.00 55.34 B ATOM 3620 CB ARG 306 8.875 -76.543 138.059 1.00 62.90 B ATOM 3622 CB ARG 306 6.644 -77.955 144.900 1.00 62.90 B ATOM 3622	5	MOTA		N	PRO	303	17.202 -74.860 137.430 1.00 74.61	
ATOM 3596 CB PRO 303 16.012 -73.215 136.261 1.00 73.72 B ROTOM 3597 CG PRO 303 15.010 -74.319 138.321 1.00 72.47 B ATOM 3599 O PRO 303 15.010 -74.319 138.321 1.00 72.47 B ATOM 3600 N ARG 304 14.920 -75.648 138.301 1.00 72.47 B ATOM 3601 CA ARG 304 12.920 -75.648 138.301 1.00 72.47 B ATOM 3601 CA ARG 304 13.796 -76.366 138.901 1.00 69.81 B ATOM 3603 CG ARG 304 13.996 -76.366 138.901 1.00 69.81 B ATOM 3603 CG ARG 304 13.996 -76.366 138.901 1.00 70.12.5 B ATOM 3603 CG ARG 304 13.994 -76.484 140.423 1.00 71.29 B ATOM 3603 CG ARG 304 14.821 -77.652 140.877 1.00 72.92 B ATOM 3605 NE ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3606 CZ ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3606 CZ ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3608 NH2 ARG 304 15.207 -79.188 142.794 1.00 76.56 B ATOM 3608 NH2 ARG 304 15.613 -80.876 144.309 1.00 76.56 B ATOM 3608 NH2 ARG 304 15.613 -80.876 144.309 1.00 76.55 B ATOM 3609 C ARG 304 11.882 -74.999 139.379 1.00 68.11 B ATOM 3610 CA ARG 304 11.882 -74.999 139.379 1.00 68.11 B ATOM 3610 CA ARG 304 11.882 -74.999 139.379 1.00 68.11 B ATOM 3613 CB ASP 305 10.708 -75.417 136.929 1.00 61.23 B ATOM 3613 CB ASP 305 10.708 -75.417 136.929 1.00 61.23 B ATOM 3615 ODI ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3616 ODI ASP 305 9.629 -77.380 135.955 1.00 62.71 B ATOM 3610 ODI ASP 305 9.629 -77.921 38.015 1.00 62.51 B ATOM 3610 CDI ASP 305 9.629 -77.921 38.015 1.00 58.07 B ATOM 3610 CDI ASP 305 9.629 -77.921 38.015 1.00 58.07 B ATOM 3610 CDI ASP 305 9.629 -77.921 38.015 1.00 58.07 B ATOM 3610 CDI ASP 305 9.629 -77.921 38.015 1.00 58.07 B ATOM 3610 CDI ASP 305 9.629 -77.981 142.941 1.00 64.52 B ATOM 3620 CDI ARG 306 8.878 7-6.531 138.016 1.00 58.07 B ATOM 3620 CDI ARG 306 9.051 -78.364 142.041 1.00 56.343 B ATOM 3620 CDI ARG 306 8.878 7-6.533 138.016 1.00 55.342 B ATOM 3620 CDI ARG 306 8.878 7-75.951 140.611 1.00 56.342 B ATOM 3620 CDI ARG 306 8.878 7-75.951 310.051 1.00 30.13 B ATOM 3630 CDI ARG 306 SOLD ARG 306 8.875 -77.381 134.4800 1.00 62.66 B ATOM 3631 CDI PHE 307 5.481 -74.0		MOTA		CD	PRO			
ATOM		MOTA		CA	PRO	303	16.273 -73.750 137.667 1.00 73.58	
10 ATOM 3599 C PRO 303 15.010 -74.319 138.321 1.00 72.47 B ATOM 3600 N ARG 304 14.920 -75.648 138.330 1.00 71.25 B ATOM 3601 CA ARG 304 14.920 -75.648 138.330 1.00 71.25 B ATOM 3601 CB ARG 304 13.976 -76.366 138.901 1.00 69.81 B ATOM 3603 CG ARG 304 13.947 -76.484 140.423 1.00 71.29 B ATOM 3604 CD ARG 304 14.821 -77.652 140.877 1.00 72.29 B ATOM 3606 CD ARG 304 14.821 -77.652 140.877 1.00 74.32 B ATOM 3606 CZ ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3606 CZ ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3606 NBL ARG 304 15.207 -79.188 142.794 1.00 75.67 B ATOM 3606 NBL ARG 304 15.207 -79.188 142.794 1.00 76.56 B ATOM 3608 NBL ARG 304 15.207 -79.188 142.794 1.00 76.56 B ATOM 3608 NBL ARG 304 15.207 -79.188 142.794 1.00 76.56 B ATOM 3610 C ARG 304 15.207 -79.188 142.794 1.00 76.56 B ATOM 3610 NBL ARG 304 15.613 -80.876 144.309 1.00 76.56 B ATOM 3610 NBL ARG 304 15.613 -80.876 144.309 1.00 76.56 B ATOM 3610 NBL ARG 304 11.882 -74.999 139.379 1.00 68.11 B ATOM 3610 C ARG 304 11.882 -74.999 139.379 1.00 68.11 B ATOM 3613 CB ASP 305 10.789 -75.956 137.344 1.00 61.23 B ATOM 3613 CB ASP 305 10.789 -75.417 136.929 1.00 61.23 B ATOM 3613 CB ASP 305 10.789 -74.232 135.882 1.00 61.23 B ATOM 3613 CB ASP 305 9.659 -74.800 135.534 1.00 61.24 B ATOM 3616 OD2 ASP 305 9.659 -74.800 135.534 1.00 61.24 B ATOM 3617 C ASP 305 9.625 -77.380 135.959 1.00 62.71 B ATOM 3617 CB ASP 305 9.625 -77.380 136.956 1.00 62.41 B ATOM 3620 CB ARG 306 8.878 -76.543 138.016 1.00 65.850 B ATOM 3621 CB ARG 306 8.878 -76.543 138.016 1.00 65.91 B ATOM 3622 CG ARG 306 8.878 -76.543 138.016 1.00 65.91 B ATOM 3622 CB ARG 306 9.051 -78.364 144.308 1.00 61.82 B ATOM 3623 CD ARG 306 8.878 -76.543 138.016 1.00 62.41 B ATOM 3625 CB ARG 306 8.878 -76.543 138.016 1.00 62.41 B ATOM 3625 CB ARG 306 9.051 -78.364 142.988 1.00 61.82 B ATOM 3625 CB ARG 306 8.878 -76.543 138.016 1.00 62.90 B ATOM 3625 CB ARG 306 8.878 -76.644 139.651 1.00 9.535 B ATOM 3626 CB ARG 306 8.878 -76.644 139.9551 1.00 9.966 B ATOM 3634 CB PHE 307 5.647 -75.583 135.807 1.00 9.966 B A		MOTA			PRO	303	16.012 -73.215 136.261 1.00 73.72	
ATOM		ATOM			PRO			
ATOM	10	ATOM			PRO		15.010 -74.319 138.321 1.00 72.47	в С
ATOM   3601   CA   ARG   304   13.996 -76.366   138.901   1.00   69.81   B   ATOM   3602   CB   ARG   304   13.947 -76.484   140.423   1.00   71.29   B   ATOM   3603   CG   ARG   304   14.821 -77.652   140.877   1.00   72.92   B   ATOM   3605   NE   ARG   304   14.673 -77.891   142.376   1.00   75.67   B   ATOM   3606   CZ   ARG   304   15.207 -79.188   142.794   1.00   75.67   B   ATOM   3606   CZ   ARG   304   15.207 -79.188   142.794   1.00   75.67   B   ATOM   3606   CZ   ARG   304   15.207 -79.188   144.907   1.00   76.55   B   ATOM   3608   NH2   ARG   304   15.103 -79.684   144.027   1.00   76.55   B   ATOM   3609   C   ARG   304   15.613 -80.876   144.309   1.00   76.34   B   ATOM   3611   N   ASP   305   11.989 -74.999   139.379   1.00   68.11   B   ATOM   3611   N   ASP   305   11.989 -75.956   137.344   1.00   64.75   B   ATOM   3612   CA   ASP   305   10.708 -75.417   136.929   1.00   61.23   B   ATOM   3614   CG   ASP   305   10.708 -74.800   135.534   1.00   61.94   B   ATOM   3616   OD2   ASP   305   9.459 -74.232   135.082   1.00   62.77   B   ATOM   3616   OD2   ASP   305   9.629 -74.232   135.082   1.00   62.77   B   ATOM   3616   OD2   ASP   305   9.629 -74.232   135.082   1.00   62.77   B   ATOM   3616   OD2   ASP   305   9.629 -74.232   135.082   1.00   62.51   B   ATOM   3616   OD2   ASP   305   9.629 -77.3803   136.956   1.00   58.07   B   ATOM   3620   CA   ARG   306   8.878 -76.543   138.10f   1.00   58.07   B   ATOM   3620   CA   ARG   306   8.878 -76.543   138.10f   1.00   58.07   B   ATOM   3620   CA   ARG   306   7.840 -77.548   138.187   1.00   51.10   B   ATOM   3622   CG   ARG   306   8.818 -77.954   142.941   1.00   62.64   B   ATOM   3625   CZ   ARG   306   8.918 -77.954   142.941   1.00   62.64   B   ATOM   3625   CZ   ARG   306   8.918 -77.954   142.941   1.00   62.64   B   ATOM   3626   NH   ARG   306   8.925 -77.3803   138.10f   1.00   59.35   B   ATOM   3626   CZ   ARG   306   6.664   -77.564   139.655   1.00   30.42   B   ATOM   3626   CZ   ARG   306   6.664   -77.564		MOTA	3599	0		303		в О
ATOM   3602   CB   ARG   304   13.947   -76.484   140.423   1.00   71.29   B   ATOM   3603   CG   ARG   304   14.821   -77.652   140.877   1.00   72.92   B   ATOM   3605   NE   ARG   304   14.673   -77.891   142.376   1.00   74.32   B   ATOM   3605   NE   ARG   304   15.207   -79.188   142.794   1.00   76.55   B   ATOM   3606   CZ   ARG   304   15.103   -79.684   144.027   1.00   76.55   B   ATOM   3608   NH2   ARG   304   15.103   -79.684   144.027   1.00   76.55   B   ATOM   3608   NH2   ARG   304   15.613   -80.876   144.980   1.00   76.55   B   ATOM   3610   O   ARG   304   12.464   -75.716   138.562   1.00   67.71   B   ATOM   3611   N   ASP   305   11.989   -75.956   137.344   1.00   64.75   B   ATOM   3611   N   ASP   305   10.708   -74.800   135.334   1.00   61.23   B   ATOM   3614   CG   ASP   305   10.708   -74.800   135.534   1.00   61.23   B   ATOM   3615   ODI   ASP   305   9.459   -74.820   135.082   1.00   62.41   B   ATOM   3617   C   ASP   305   9.459   -74.820   135.082   1.00   62.41   B   ATOM   3618   ODI   ASP   305   9.679   -76.536   136.956   1.00   62.51   B   ATOM   3618   ODI   ASP   305   9.679   -76.536   136.956   1.00   62.51   B   ATOM   3618   ODI   ASP   305   9.679   -76.536   136.956   1.00   58.50   B   ATOM   3618   ODI   ASP   305   9.679   -76.536   136.956   1.00   58.50   B   ATOM   3618   ODI   ASP   305   9.679   -76.536   136.956   1.00   58.50   B   ATOM   3620   CDI   ASP   305   9.679   -76.536   136.956   1.00   58.50   B   ATOM   3621   CDI   ARG   306   8.878   -77.891   142.041   1.00   59.35   B   ATOM   3622   CDI   ARG   306   8.878   -77.548   138.016   1.00   54.93   B   ATOM   3622   CDI   ARG   306   8.878   -77.548   138.016   1.00   54.93   B   ATOM   3622   CDI   ARG   306   8.878   -77.588   142.041   1.00   59.35   B   ATOM   3622   CDI   ARG   306   8.878   -77.588   144.308   1.00   61.82   B   ATOM   3628   CDI   ARG   306   6.644   -77.798   144.308   1.00   61.62   B   ATOM   3628   CDI   ARG   306   6.644   -77.986   144.308   1.00   61.62		MOTA	3600	N	ARG	304	14.920 -75.648 138.330 1.00 71.25	B N
15		MOTA	3601	CA	ARG	304		в С
ATOM   3604   CD   ARG   304   14.673   -77.891   142.376   1.00   74.32   B   ATOM   3605   NE   ARG   304   15.207   -79.188   142.376   1.00   75.67   B   ATOM   3607   NH1   ARG   304   15.207   -79.684   144.027   1.00   76.25   B   ATOM   3608   NH2   ARG   304   14.484   -78.995   144.980   1.00   76.56   B   ATOM   3608   NH2   ARG   304   15.613   -80.876   144.399   1.00   76.34   B   ATOM   3610   O   ARG   304   11.882   -74.999   139.379   1.00   67.71   B   ATOM   3611   N   ASP   305   11.989   -75.956   137.344   1.00   64.75   B   ATOM   3612   CA   ASP   305   10.708   -75.956   137.344   1.00   64.75   B   ATOM   3613   CB   ASP   305   10.708   -75.401   136.929   1.00   61.23   B   ATOM   3614   CG   ASP   305   9.459   -74.800   135.534   1.00   62.41   B   ATOM   3615   ODI   ASP   305   8.677   -73.803   135.959   1.00   62.41   B   ATOM   3616   OD2   ASP   305   9.627   -73.803   135.959   1.00   62.51   B   ATOM   3617   C   ASP   305   9.627   -74.194   133.860   1.00   58.50   B   ATOM   3618   O   ASP   305   9.627   -74.194   133.860   1.00   58.50   B   ATOM   3619   N   ARG   306   8.878   -76.543   138.016   1.00   54.93   B   ATOM   3620   CB   ARG   306   7.480   -77.548   138.107   1.00   54.93   B   ATOM   3622   CB   ARG   306   7.480   -77.595   140.611   1.00   56.34   B   ATOM   3622   CB   ARG   306   7.480   -77.595   140.611   1.00   56.34   B   ATOM   3625   CB   ARG   306   7.480   -77.595   140.611   1.00   56.34   B   ATOM   3625   CB   ARG   306   8.513   -77.985   140.611   1.00   56.34   B   ATOM   3625   CB   ARG   306   7.480   -77.595   140.611   1.00   56.34   B   ATOM   3625   CB   ARG   306   6.644   -77.595   140.611   1.00   56.34   B   ATOM   3625   CB   ARG   306   6.644   -77.924   137.318   1.00   61.82   B   ATOM   3625   CB   ARG   306   6.644   -77.595   140.611   1.00   56.34   B   ATOM   3626   NH1   ARG   306   6.644   -77.595   140.611   1.00   36.02   B   ATOM   3636   CB   HE   307   5.6481   -77.955   315.602   1.00   38.24   B   ATOM		MOTA	3602	CB	ARG	304	13.947 -76.484 140.423 1.00 71.29	в С
ATOM   3605   NE   ARG   304   15.207 -79.188   142.794   1.00   75.67   B   ATOM   3606   CZ   ARG   304   15.103 -79.684   144.027   1.00   76.25   B   ATOM   3608   NH1   ARG   304   14.484   -78.995   144.980   1.00   76.56   B   ATOM   3608   NH2   ARG   304   15.613   -80.876   144.309   1.00   76.34   B   ATOM   3610   O   ARG   304   11.882   -74.999   139.379   1.00   68.11   B   ATOM   3611   N   ASP   305   11.989   -75.956   137.344   1.00   64.75   B   ATOM   3612   CA   ASP   305   10.789   -74.800   135.534   1.00   61.23   B   ATOM   3615   ODI   ASP   305   10.789   -74.800   135.534   1.00   61.23   B   ATOM   3615   ODI   ASP   305   8.677   -73.803   135.959   1.00   62.41   B   ATOM   3616   ODI   ASP   305   8.677   -73.803   135.959   1.00   62.41   B   ATOM   3616   ODI   ASP   305   9.679   -76.536   136.956   1.00   62.51   B   ATOM   3618   O   ASP   305   9.679   -76.536   136.956   1.00   62.51   B   ATOM   3618   O   ASP   305   9.679   -76.536   136.956   1.00   58.50   B   ATOM   3618   O   ASP   305   9.679   -76.536   136.956   1.00   54.93   B   ATOM   3620   CA   ARG   306   R.878   -77.3803   138.916   1.00   54.93   B   ATOM   3620   CA   ARG   306   7.840   -77.584   138.187   1.00   51.10   B   ATOM   3622   CB   ARG   306   8.016   -77.955   140.611   1.00   59.35   B   ATOM   3623   CD   ARG   306   8.016   -77.955   140.611   1.00   59.35   B   ATOM   3623   CD   ARG   306   8.016   -77.955   140.611   1.00   59.35   B   ATOM   3625   CD   ARG   306   8.016   -77.955   140.611   1.00   50.34   B   ATOM   3626   CD   ARG   306   8.016   -77.955   140.611   1.00   50.34   B   ATOM   3626   CD   ARG   306   8.016   -77.955   140.611   1.00   50.34   B   ATOM   3628   CD   ARG   306   8.016   -77.955   140.611   1.00   50.34   B   ATOM   3626   CD   ARG   306   8.016   -77.955   140.611   1.00   50.34   B   ATOM   3630   CD   PHE   307   5.681   -77.955   140.611   1.00   40.629   B   ATOM   3636   CD   PHE   307   5.681   -77.955   137.341   1.00   30.966   B   ATOM   3	15	MOTA	3603	CG	ARG	304	14.821 -77.652 140.877 1.00 72.92	в С
ATOM		ATOM	3604	CD	ARG	304	14.673 -77.891 142.376 1.00 74.32	в С
ATOM   3607   NH1   ARG   304   14.484   -78.995   144.980   1.00   76.56   8		MOTA	3605	NE	ARG	304	15.207 -79.188 142.794 1.00 75.67	B N
200		MOTA	3606	CZ	ARG	304	15.103 -79.684 144.027 1.00 76.25	в с
ATOM 3610 C ARG 304 12.464 -75.716 138.562 1.00 67.71 B ATOM 3610 O ARG 304 11.882 -74.999 139.379 1.00 68.11 B ATOM 3611 N ASP 305 11.989 -75.956 137.341 1.00 64.75 B ATOM 3612 CA ASP 305 10.708 -75.417 136.929 1.00 61.23 B ATOM 3613 CB ASP 305 10.789 -74.800 135.534 1.00 61.24 B ATOM 3613 CB ASP 305 10.789 -74.800 135.534 1.00 61.94 B ATOM 3615 OD1 ASP 305 9.459 -74.232 135.082 1.00 62.41 B ATOM 3616 OD2 ASP 305 9.202 -74.194 133.860 1.00 62.77 B ATOM 3617 C ASP 305 9.202 -74.194 133.860 1.00 62.51 B ATOM 3618 O ASP 305 9.202 -74.194 133.860 1.00 62.51 B ATOM 3618 O ASP 305 9.679 -76.536 136.956 1.00 58.50 B ATOM 3618 O ASP 305 9.625 -77.382 136.059 1.00 58.07 B ATOM 3620 CA ARG 306 8.878 -76.543 138.016 1.00 54.93 B ATOM 3620 CA ARG 306 7.400 -77.548 138.106 1.00 54.93 B ATOM 3620 CA ARG 306 7.400 -77.604 139.651 1.00 55.42 B ATOM 3623 CD ARG 306 8.516 -77.985 140.611 1.00 56.34 B ATOM 3623 CD ARG 306 8.516 -77.985 140.611 1.00 56.34 B ATOM 3623 CD ARG 306 8.516 -77.985 140.611 1.00 56.34 B ATOM 3626 NH1 ARG 306 8.925 -78.298 144.308 1.00 62.64 B ATOM 3626 NH1 ARG 306 9.051 -78.364 142.988 1.00 61.82 B ATOM 3626 NH1 ARG 306 9.051 -78.364 142.988 1.00 61.82 B ATOM 3630 N PHE 307 6.644 -77.204 137.318 1.00 62.64 B ATOM 3630 N PHE 307 6.644 -77.204 137.318 1.00 62.64 B ATOM 3631 CA PHE 307 5.647 -75.583 135.807 1.00 38.24 B ATOM 3632 CD PHE 307 5.647 -75.583 135.807 1.00 38.24 B ATOM 3632 CD PHE 307 5.647 -75.583 135.807 1.00 38.24 B ATOM 3633 CD PHE 307 5.647 -75.583 135.807 1.00 38.24 B ATOM 3636 CD1 PHE 307 5.647 -75.583 139.047 1.00 36.42 B ATOM 3637 CD PHE 307 5.915 -73.160 139.933 1.00 40.42 B ATOM 3637 CD PHE 307 5.915 -73.160 139.933 1.00 40.42 B ATOM 3634 CD PHE 307 5.915 -73.160 139.933 1.00 40.42 B ATOM 3634 CD PHE 307 5.915 -73.160 139.933 1.00 40.42 B ATOM 3634 CD PHE 307 5.915 -73.160 139.933 1.00 40.42 B ATOM 3636 CD PHE 307 5.915 -73.160 139.933 1.00 40.42 B ATOM 3636 CD PHE 307 5.915 -73.160 139.933 1.00 40.42 B ATOM 3634 CD PHE 307 5.905 -75.711 133.494 1.00 30.13 B ATOM 3645 CD PHE 307 5.905		ATOM	3607	NH1	ARG	304	14.484 -78.995 144.980 1.00 76.56	B N
ATOM   3610   O   ARG   304	20	ATOM	3608	NH2	ARG	304	15.613 -80.876 144.309 1.00 76.34	B N
ATOM		MOTA	3609	С	ARG	304	12.464 -75.716 138.562 1.00 67.71	в с
ATOM		ATOM	3610	0	ARG	304	11.882 -74.999 139.379 1.00 68.11	в о
25         ATOM         3613         CB         ASP         305         10.789         -74.800         135.534         1.00         61.94         B           ATOM         3615         ODI         ASP         305         9.459         -74.232         135.082         1.00         62.41         B           ATOM         3616         ODZ         ASP         305         9.202         -74.194         133.860         1.00         62.51         B           ATOM         3616         ODZ         ASP         305         9.679         -76.536         136.059         1.00         58.50         B           ATOM         3618         O         ASP         305         9.625         -77.382         136.059         1.00         58.07         B           ATOM         3620         CA         ARG         306         8.878         -76.543         138.016         1.00         53.42         B           ATOM         3621         CB         ARG         306         7.408         -77.548         138.187         1.00         53.42         B           ATOM         3621         CB         ARG         306         8.513         -77.954         140.611		MOTA	3611	N	ASP	305	11.989 -75.956 137.344 1.00 64.75	B N
ATOM 3614 CG ASP 305 9.459 -74.232 135.082 1.00 62.41 B ATOM 3615 OD1 ASP 305 8.677 -73.803 135.952 1.00 62.77 B ATOM 3616 OD2 ASP 305 9.202 -74.194 133.860 1.00 62.51 B ATOM 3617 C ASP 305 9.679 -76.536 136.956 1.00 58.50 B ATOM 3618 O ASP 305 9.625 -77.382 136.059 1.00 58.07 B ATOM 3619 N ARG 306 8.878 -76.543 138.016 1.00 54.93 B ATOM 3620 CA ARG 306 7.840 -77.548 138.187 1.00 51.10 B ATOM 3621 CB ARG 306 7.408 -77.604 139.651 1.00 53.42 B ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3623 CD ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3624 NE ARG 306 8.016 -77.954 140.441 1.00 59.35 B ATOM 3625 CZ ARG 306 8.925 -78.298 144.308 1.00 61.82 B ATOM 3626 NH1 ARG 306 8.925 -78.298 144.308 1.00 62.64 B ATOM 3627 NH2 ARG 306 7.805 -77.831 144.840 1.00 63.33 B ATOM 3628 C ARG 306 6.644 -77.204 137.318 1.00 46.82 B ATOM 3629 O ARG 306 5.683 -77.962 137.241 1.00 46.07 B ATOM 3631 CA PHE 307 5.647 -75.583 135.807 1.00 38.22 B ATOM 3632 CB PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3633 CG PHE 307 5.481 -74.068 135.962 1.00 38.97 B ATOM 3636 CE1 PHE 307 5.481 -74.068 135.962 1.00 38.97 B ATOM 3637 CC2 PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3631 CA PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3636 CE1 PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3637 CE2 PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3631 CB PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3634 CD1 PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3636 CE1 PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3638 CZ PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3630 C PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3640 C PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3641 N LEU 308 7.085 -76.445 134.036 1.00 32.23 B ATOM 3644 CG LEU 308 8.818 -77.445 132.667 1.00 30.13 B ATOM 3644 CG LEU 308 8.818 -77		MOTA	3612	CA	ASP	305	10.708 -75.417 136.929 1.00 61.23	в с
ATOM 3615 OD1 ASP 305 8.677 -73.803 135.959 1.00 62.77 B ATOM 3616 OD2 ASP 305 9.202 -74.194 133.860 1.00 62.51 B ATOM 3617 C ASP 305 9.679 -76.536 136.956 1.00 58.50 B ATOM 3618 O ASP 305 9.625 -77.382 136.059 1.00 58.07 B ATOM 3619 N ARG 306 8.878 -76.543 138.016 1.00 54.93 B ATOM 3620 CA ARG 306 7.840 -77.548 138.196 1.00 51.10 B ATOM 3621 CB ARG 306 7.408 -77.604 139.651 1.00 55.34 B ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3622 CG ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3624 NE ARG 306 9.051 -78.364 142.988 1.00 61.82 B ATOM 3625 CZ ARG 306 8.925 -78.298 144.308 1.00 62.64 B ATOM 3626 NH1 ARG 306 9.923 -78.686 145.090 1.00 62.90 B ATOM 3627 NH2 ARG 306 7.805 -77.831 144.840 1.00 63.33 B ATOM 3628 C ARG 306 6.644 -77.204 137.318 1.00 46.07 B ATOM 3630 N PHE 307 6.714 -76.049 136.668 1.00 42.30 B ATOM 3631 CA PHE 307 5.647 -75.583 135.807 1.00 38.22 B ATOM 3632 CB PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3635 CD2 PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3636 CE1 PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3636 CE1 PHE 307 5.179 -73.628 137.369 1.00 38.97 B ATOM 3636 CE2 PHE 307 5.179 -73.628 137.369 1.00 38.97 B ATOM 3636 CE1 PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3637 CE2 PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3639 C PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3640 O PHE 307 5.901 -75.924 134.336 1.00 32.23 B ATOM 3640 CD PHE 307 5.901 -75.924 134.336 1.00 32.23 B ATOM 3641 N LEU 308 7.085 -76.445 134.036 1.00 32.23 B ATOM 3642 CA LEU 308 8.818 -77.445 132.667 1.00 30.71 B ATOM 3644 CG LEU 308 8.818 -77.445 132.667 1.00 30.71 B ATOM 3644 CG LEU 308 8.97.5 -77.108 131.347 1.00 30.71 B ATOM 3645 CD1 LEU 308 8.97.5 -77.108 131.314 1.00 31.09 B	25	MOTA	3613	CB	ASP	305	10.789 -74.800 135.534 1.00 61.94	в с
ATOM 3616 OD2 ASP 305 9.202 -74.194 133.860 1.00 62.51 B ATOM 3617 C ASP 305 9.679 -76.536 136.956 1.00 58.50 B ATOM 3618 O ASP 305 9.625 -77.382 136.059 1.00 58.07 B ATOM 3619 N ARG 306 8.878 -76.543 138.016 1.00 54.93 B ATOM 3620 CA ARG 306 7.840 -77.548 138.187 1.00 51.10 B ATOM 3621 CB ARG 306 7.408 -77.604 139.651 1.00 53.42 B ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 59.35 B ATOM 3623 CD ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3624 NE ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3625 CZ ARG 306 8.925 -78.298 144.308 1.00 61.82 B ATOM 3626 NH1 ARG 306 9.051 -78.364 142.988 1.00 62.64 B ATOM 3626 NH1 ARG 306 9.923 -78.686 145.090 1.00 62.90 B ATOM 3626 NH2 ARG 306 6.644 -77.204 137.328 1.00 46.82 B ATOM 3629 O ARG 306 6.644 -77.204 137.328 1.00 46.82 B ATOM 3630 N PHE 307 6.714 -76.049 136.668 1.00 42.30 B ATOM 3631 CA PHE 307 5.481 -74.068 135.962 1.00 38.22 B ATOM 3632 CB PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3633 CG PHE 307 5.179 -73.628 137.369 1.00 38.24 B ATOM 3636 CE1 PHE 307 5.179 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.179 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.179 -73.628 137.369 1.00 39.66 B ATOM 3636 CE2 PHE 307 5.179 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3637 CE2 PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3638 CZ PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3639 C PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3630 CE1 PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3630 CE1 PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3630 CE1 PHE 307 5.901 -75.924 134.337 1.00 32.23 B ATOM 3640 C PHE 307 5.901 -75.924 134.337 1.00 32.23 B ATOM 3642 CG LEU 308 7.441 -76.776 132.663 1.00 39.80 B ATOM 3644 CG LEU 308 7.441 -76.776 132.663 1.00 39.60 B ATOM 3644 CG LEU 308 8.818 -77.445 132.663 1.00 39.01 B ATOM 3644 CG LEU 308 8.818 -77.445 132.661 1.00 30.71 B ATOM 3645 CD1 LEU 308 7.257 -77.108 131.314 1.00 31.09 B		ATOM	3614	CG	ASP	305	9.459 -74.232 135.082 1.00 62.41	в с
ATOM 3618 O ASP 305 9.679 -76.536 136.956 1.00 58.50 B ATOM 3618 O ASP 305 9.625 -77.382 136.059 1.00 58.07 B ATOM 3619 N ARG 306 8.878 -76.543 138.016 1.00 54.93 B ATOM 3620 CA ARG 306 7.840 -77.548 138.187 1.00 51.10 B ATOM 3621 CB ARG 306 7.408 -77.604 139.651 1.00 53.42 B ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3623 CD ARG 306 8.513 -77.985 140.611 1.00 59.35 B ATOM 3624 NE ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3625 CZ ARG 306 8.925 -78.298 144.308 1.00 61.82 B ATOM 3626 NH1 ARG 306 9.051 -78.364 142.988 1.00 62.64 B ATOM 3627 NH2 ARG 306 7.805 -77.831 144.840 1.00 63.33 B ATOM 3628 C ARG 306 7.805 -77.831 144.840 1.00 63.33 B ATOM 3629 O ARG 306 5.683 -77.962 137.241 1.00 46.07 B ATOM 3630 N PHE 307 6.714 -76.049 136.668 1.00 42.30 B ATOM 3631 CA PHE 307 5.647 -75.583 135.962 1.00 38.22 B ATOM 3632 CB PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3633 CG PHE 307 5.481 -74.068 135.962 1.00 38.97 B ATOM 3636 CE1 PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3638 CZ PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3639 C PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3630 N PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3637 CE2 PHE 307 5.915 -73.628 137.369 1.00 39.66 B ATOM 3638 CZ PHE 307 5.915 -73.628 133.993 1.00 40.42 B ATOM 3630 N PHE 307 5.915 -73.628 133.993 1.00 40.42 B ATOM 3630 N PHE 307 5.915 -73.160 139.632 1.00 39.66 B ATOM 3630 CE1 PHE 307 5.915 -77.181 133.494 1.00 34.63 B ATOM 3630 CE1 PHE 307 5.915 -77.485 134.036 1.00 39.22 B ATOM 3640 CE1 PHE 307 5.030 -75.731 133.494 1.00 34.63 B ATOM 3640 CE LEU 308 8.818 -77.445 132.667 1.00 30.13 B ATOM 3644 CG LEU 308 8.818 -77.445 132.617 1.00 30.71 B ATOM 3645 CD1 LEU 308 8.818 -77.445 132.617 1.00 30.71 B		ATOM	3615	OD1	ASP	305	8.677 -73.803 135.959 1.00 62.77	в о
ATOM   3618   O   ASP   305   9.625   -77.382   136.059   1.00   58.07   B		MOTA	3616	OD2	ASP	305	9.202 -74.194 133.860 1.00 62.51	в о
ATOM 3619 N ARG 306 8.878 -76.543 138.016 1.00 54.93 B ATOM 3620 CA ARG 306 7.840 -77.548 138.187 1.00 51.10 B ATOM 3621 CB ARG 306 7.408 -77.604 139.651 1.00 53.42 B ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3623 CD ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3624 NE ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3625 CZ ARG 306 8.925 -78.298 144.308 1.00 62.64 B ATOM 3626 NH1 ARG 306 8.925 -78.298 144.308 1.00 62.64 B ATOM 3626 NH1 ARG 306 9.923 -78.686 145.090 1.00 62.09 B ATOM 3627 NH2 ARG 306 7.805 -77.831 144.840 1.00 63.33 B ATOM 3629 O ARG 306 6.644 -77.204 137.318 1.00 46.07 B ATOM 3630 N PHE 307 6.714 -76.049 136.668 1.00 42.30 B ATOM 3631 CA PHE 307 5.647 -75.583 135.807 1.00 38.22 B ATOM 3631 CA PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3633 CG PHE 307 5.481 -74.068 135.962 1.00 38.97 B ATOM 3636 CD2 PHE 307 5.179 -73.628 137.369 1.00 38.97 B ATOM 3636 CE1 PHE 307 3.883 -73.556 138.326 1.00 39.66 B ATOM 3637 CE2 PHE 307 3.883 -73.596 137.741 1.00 30.66 B ATOM 3638 CZ PHE 307 3.883 -73.296 137.741 1.00 39.64 B ATOM 3638 CZ PHE 307 3.883 -73.296 137.741 1.00 39.64 B ATOM 3639 C PHE 307 5.915 -73.160 139.933 1.00 40.42 B ATOM 3639 C PHE 307 5.901 -75.924 134.337 1.00 32.23 B ATOM 3639 C PHE 307 5.901 -75.924 134.337 1.00 32.23 B ATOM 3639 C PHE 307 5.901 -75.924 134.337 1.00 32.23 B ATOM 3639 C PHE 307 5.901 -75.924 134.337 1.00 32.23 B ATOM 3640 O PHE 307 5.901 -75.924 134.337 1.00 32.23 B ATOM 3640 O PHE 307 5.901 -75.924 134.337 1.00 32.23 B ATOM 3641 N LEU 308 7.441 -76.776 132.663 1.00 29.80 B ATOM 3643 CB LEU 308 7.441 -76.776 132.663 1.00 29.80 B ATOM 3644 CG LEU 308 9.725 -77.108 131.344 1.00 31.09 B		ATOM	3617	С	ASP	305	9.679 -76.536 136.956 1.00 58.50	в с
ATOM 3620 CA ARG 306 7.840 -77.548 138.187 1.00 51.10 B ATOM 3621 CB ARG 306 7.408 -77.604 139.651 1.00 53.42 B ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3623 CD ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3624 NE ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3624 NE ARG 306 9.051 -78.364 142.988 1.00 61.82 B ATOM 3625 CZ ARG 306 8.925 -78.298 144.308 1.00 62.64 B ATOM 3626 NH1 ARG 306 9.923 -78.686 145.090 1.00 62.90 B ATOM 3627 NH2 ARG 306 7.805 -77.831 144.840 1.00 63.33 B ATOM 3628 C ARG 306 6.644 -77.204 137.318 1.00 46.82 B ATOM 3629 O ARG 306 6.644 -77.204 137.318 1.00 46.82 B ATOM 3630 N PHE 307 6.714 -76.049 136.668 1.00 42.30 B ATOM 3631 CA PHE 307 5.647 -75.583 135.807 1.00 38.22 B ATOM 3633 CG PHE 307 5.481 -74.668 135.962 1.00 38.24 B ATOM 3633 CG PHE 307 5.179 -73.628 137.369 1.00 38.24 B ATOM 3634 CD1 PHE 307 6.188 -73.556 138.326 1.00 39.66 B ATOM 3635 CD2 PHE 307 3.883 -73.296 137.741 1.00 39.64 B ATOM 3636 CE1 PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3638 CD PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3638 CZ PHE 307 5.915 -75.924 134.337 1.00 35.22 B ATOM 3638 CZ PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3634 CB LEU 308 7.085 -76.445 134.036 1.00 39.63 B ATOM 3641 N LEU 308 7.085 -76.445 134.036 1.00 39.80 B ATOM 3644 CG LEU 308 8.818 -77.445 132.617 1.00 30.11 B ATOM 3644 CG LEU 308 8.818 -77.445 132.617 1.00 30.11 B ATOM 3644 CG LEU 308 8.818 -77.445 132.617 1.00 30.11 B ATOM 3644 CG LEU 308 8.818 -77.445 132.617 1.00 30.11 B ATOM 3644 CG LEU 308 9.725 -77.108 131.314 1.00 31.09 B	30	ATOM	3618	0	ASP	305	9.625 -77.382 136.059 1.00 58.07	в о
ATOM 3621 CB ARG 306 7.408 -77.604 139.651 1.00 53.42 B ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3623 CD ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3624 NE ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3625 CZ ARG 306 8.925 -78.364 142.988 1.00 61.82 B ATOM 3626 NH1 ARG 306 8.925 -78.298 144.308 1.00 62.64 B ATOM 3627 NH2 ARG 306 9.923 -78.686 145.090 1.00 62.90 B ATOM 3628 C ARG 306 7.805 -77.831 144.840 1.00 63.33 B ATOM 3629 O ARG 306 6.644 -77.204 137.318 1.00 46.82 B ATOM 3630 N PHE 307 6.714 -76.049 136.668 1.00 42.30 B ATOM 3631 CA PHE 307 5.647 -75.583 135.807 1.00 38.22 B ATOM 3633 CG PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3633 CG PHE 307 5.179 -73.628 137.369 1.00 38.97 B ATOM 3634 CD1 PHE 307 6.188 -73.556 138.326 1.00 39.66 B ATOM 3635 CD2 PHE 307 3.883 -73.296 137.741 1.00 39.66 B ATOM 3636 CE1 PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3639 C PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3634 CB LEU 308 7.085 -76.445 134.036 1.00 32.23 B ATOM 3640 C PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3641 N LEU 308 7.085 -76.445 134.036 1.00 32.23 B ATOM 3644 CG LEU 308 8.818 -77.445 132.617 1.00 30.71 B ATOM 3644 CG LEU 308 9.725 -77.108 131.427 1.00 30.71 B ATOM 3644 CG LEU 308 9.725 -77.108 131.427 1.00 30.71 B ATOM 3644 CG LEU 308 9.725 -77.108 131.427 1.00 30.71 B		MOTA	3619	N	ARG	306	8.878 -76.543 138.016 1.00 54.93	B N
ATOM 3622 CG ARG 306 8.513 -77.985 140.611 1.00 56.34 B ATOM 3624 NE ARG 306 8.016 -77.954 142.041 1.00 59.35 B ATOM 3624 NE ARG 306 9.051 -78.364 142.988 1.00 61.82 B ATOM 3625 CZ ARG 306 8.925 -78.298 144.308 1.00 62.64 B ATOM 3626 NH1 ARG 306 9.923 -78.686 145.090 1.00 62.90 B ATOM 3627 NH2 ARG 306 7.805 -77.831 144.840 1.00 63.33 B ATOM 3628 C ARG 306 6.644 -77.204 137.318 1.00 46.82 B ATOM 3630 N PHE 307 6.714 -76.049 136.668 1.00 42.30 B ATOM 3631 CA PHE 307 5.647 -75.583 135.807 1.00 38.22 B ATOM 3632 CB PHE 307 5.481 -74.068 135.962 1.00 38.24 B ATOM 3634 CD1 PHE 307 5.179 -73.628 137.369 1.00 38.24 B ATOM 3635 CD2 PHE 307 5.179 -73.628 137.369 1.00 39.66 B ATOM 3636 CE1 PHE 307 3.883 -73.296 137.741 1.00 40.08 B ATOM 3636 CE2 PHE 307 5.915 -73.160 139.632 1.00 40.08 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.632 1.00 40.42 B ATOM 3638 CZ PHE 307 5.915 -73.160 139.993 1.00 40.42 B ATOM 3638 CZ PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3638 CZ PHE 307 5.901 -75.924 134.337 1.00 35.22 B ATOM 3636 CE1 PHE 307 5.030 -75.731 133.494 1.00 34.63 B ATOM 3636 CE PHE 307 5.030 -75.731 133.494 1.00 34.63 B ATOM 3636 CB LEU 308 7.085 -76.445 134.036 1.00 29.80 B ATOM 3640 CB LEU 308 7.441 -76.76 132.663 1.00 29.80 B ATOM 3644 CG LEU 308 7.441 -76.76 132.661 1.00 30.71 B ATOM 3644 CG LEU 308 9.725 -77.108 131.427 1.00 30.71 B ATOM 3644 CG LEU 308 9.725 -77.108 131.427 1.00 30.71 B ATOM 3644 CG LEU 308 9.725 -77.108 131.427 1.00 30.71 B		ATOM	3620	CA	ARG	306	7.840 -77.548 138.187 1.00 51.10	в с
35         ATOM         3623         CD         ARG         306         8.016         -77.954         142.041         1.00         59.35         B           ATOM         3624         NE         ARG         306         9.051         -78.364         142.988         1.00         61.82         B           ATOM         3625         CZ         ARG         306         8.925         -78.298         144.308         1.00         62.64         B           ATOM         3626         NH1         ARG         306         9.923         -78.686         145.090         1.00         62.90         B           ATOM         3627         NH2         ARG         306         7.805         -77.831         144.840         1.00         62.90         B           ATOM         3628         C         ARG         306         6.644         -77.204         137.318         1.00         46.82         B           ATOM         3631         CA         PHE         307         6.714         -76.049         136.668         1.00         42.30         B           ATOM         3631         CA         PHE         307         5.481         -74.068         135.962		ATOM	3621	CB	ARG	306	7.408 -77.604 139.651 1.00 53.42	вс
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ATOM 3646 CD2 LEU 308 8.932 -77.041 130.142 1.00 30.29 B								_
		MOTA	3646	CD2	LEU	308	8.932 -77.041 130.142 1.00 30.29	в с

	MOTA	3647	С	LEU	308		-77.671		1.00 27.96	В	С
	MOTA	3648	0	LEU	308		-77.296		1.00 26.38	В	0
	MOTA	3649	N	TYR	309		-78.847		1.00 26.33	В	N
_	MOTA	3650	CA	TYR	309		-79.770		1.00 26.08	В	С
5	MOTA	3651	CB	TYR	309		-81.040		1.00 26.27	В	C
	MOTA	3652	CG	TYR	309		-82.074		1.00 26.57	В	С
	MOTA	3653	CD1		309		-82.611		1.00 26.55	В	С
	MOTA	3654	CE1		309		-83.533		1.00 27.76	В	С
40	ATOM	3655	CD2		309		-82.489		1.00 26.42	В	С
10	MOTA	3656	CE2	TYR	309		-83.414		1.00 28.03	В	С
	MOTA	3657	CZ	TYR	309	2.404	-83.930	130.902	1.00 27.91	В	С
	ATOM	3658	OH	TYR	309		-84.847		1.00 29.13	В	0
	MOTA	3659	С	TYR	309		-79.135		1.00 25.30	В	C
	MOTA	3660	0	TYR	309		-79.280		1.00 24.69	В	0
15	MOTA	3661	N	ALA	310			132.649	1.00 24.77	В	N
	ATOM	3662	CA	ALA	310	2.007	-77.780	132.519	1.00 23.64	В	C
	MOTA	3663	CB	ALA	310	1.628	-77.092	133.822	1.00 24.37	В	С
	ATOM	3664	С	ALA	310	2.047	-76.764	131.385	1.00 23.22	В	С
	MOTA	3665	0	ALA	310	1.088	-76.630	130.628	1.00 22.01	В	0
20	MOTA	3666	N	LYS	311	3.158	-76.043	131.276	1.00 22.47	В	N
	MOTA	3667	CA	LYS	311	3.315	-75.052	130.217	1.00 22.76	В	С
	MOTA	3668	CB	LYS	311	4.612	-74.271	130.413	1.00 23.73	В	С
	MOTA	3669	CG	LYS	311	4.563	-73.270	131.550	1.00 25.88	В	С
	MOTA	3670	CD	LYS	311	5.880	-72.533	131.657	1.00 27.25	В	С
25	MOTA	3671	CE	LYS	311			132.820	1.00 29.50	В	С
	ATOM	3672	NZ	LYS	311			133.000	1.00 31.45	В	N
	ATOM	3673	С	LYS	311	3.309	-75.710	128.838	1.00 20.98	В	C
	ATOM	3674	0	LYS	311			127.900	1.00 21.17	В	ō
	ATOM	3675	N	LEU	312			128.721	1.00 20.58	В	N
30	ATOM	3676	CA	LEU	312			127.454	1.00 20.47	В	C
-	ATOM	3677	СВ	LEU	312			127.550	1.00 20.31	В	Č
	ATOM	3678	CG	LEU	312			127.715	1.00 20.12	В	č
	ATOM	3679		LEU	312			127.750	1.00 20.29	В	c
	ATOM	3680		LEU	312			126.561	1.00 19.66	В	Č
35	ATOM	3681	C	LEU	312			127.029	1.00 20.37	В	Č
	ATOM	3682	ō	LEU	312			125.840	1.00 20.37	В	õ
	ATOM	3683	N	LEU	313			127.989	1.00 19.73	В	Ŋ
	ATOM	3684	CA	LEU	313			127.659	1.00 19.16	В	C
	MOTA	3685	CB	LEU	313			128.872	1.00 18.30	В	c
40	ATOM	3686	CG	LEU	313			129.369	1.00 10.30	В	C
	ATOM	3687		LEU	313			130.417	1.00 13.24	В	C
	ATOM	3688		LEU	313			128.205	1.00 10.74	В	
	ATOM	3689	C	LEU	313			127.173	1.00 19.01	В	C
	ATOM	3690	ŏ	LEU	313			126.235	1.00 18.15	В	0
45	ATOM	3691	N	GLY	314			127.810	1.00 18.13	В	
70	ATOM	3692	CA	GLY	314			127.010	1.00 19.37	В	N
	ATOM	3693	C	GLY	314			127.402	1.00 19.37		C
	ATOM	3694	0	GLY	314			125.208		В	C
	ATOM	3695	N	LEU	315			125.208	1.00 20.34	В	0
50	ATOM	3696	CA	LEU	315				1.00 20.42	В	N
30		3697	CB					124.367	1.00 20.97	В	C
	MOTA			LEU	315			124.368	1.00 20.14	В	C
	ATOM	3698	CG CD1	LEU	315			125.027	1.00 21.61	В	C
	ATOM	3699		LEU	315			125.204	1.00 22.43	В	C
55	MOTA	3700		LEU	315			124.174	1.00 20.16	В	C
33	MOTA	3701	C	LEU	315			123.318	1.00 20.93	В	С
	ATOM	3702	0	LEU	315			122.200	1.00 21.86	В	0
	MOTA	3703	N	LEU	316			123.670	1.00 20.78	В	N
	MOTA	3704	CA	LEU	316	U.140	-78.038	122.728	1.00 22.07	В	С

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	ATOM	3705	СВ	LEU	316	0.122 -	79.442	123.333	1.00 22.61	В	С
	ATOM	3706	CG	LEU	316	1.463 -	80.171	123.415	1.00 25.13	В	С
	ATOM	3707	CD1		316			123.994	1.00 25.21	В	С
	ATOM	3708	CD2		316			122.021	1.00 27.32	В	C
5	ATOM	3709	C	LEU	316	-1.286 -			1.00 22.16	В	C
	ATOM	3710	ō	LEU	316	-1.723 -			1.00 21.15	В	ō
	MOTA	3711	N	ALA	317	-1.998 -			1.00 22.21	В	N
	ATOM	3712	CA	ALA	317			123.123	1.00 22.21	В	C
		3713	CB	ALA	317			124.482	1.00 22.08		c
10	ATOM							122.284		В	
10	ATOM	3714	C	ALA	317				1.00 23.40	В	C
	MOTA	3715	0	ALA	317			121.437	1.00 22.48	В	0
	MOTA	3716	N	GLU	318			122.527	1.00 23.64	В	N
	MOTA	3717	CA	GLU	318			121.763	1.00 25.42	В	C
4-	MOTA	3718	СВ	GLU	318			122.325	1.00 27.16	В	С
15	MOTA	3719	CG	GLU	318			121.859	1.00 32.13	В	С
	MOTA	3720	CD	GLU	318			122.233	1.00 34.93	В	С
	ATOM	3721		GLU	318			123.362	1.00 37.73	В	0
	MOTA	3722	OE2	GLU	318	0.239	-69.114	121.402	1.00 36.75	В	0
	MOTA	3723	С	GLU	318	-2.036	-73.490	120.298	1.00 25.14	В	С
20	ATOM	3724	0	GLU	318	-2.557	~72.851	119.384	1.00 23.94	В	0
	MOTA	3725	N	LEU	319	-1.186	-74.490	120.084	1.00 25.66	В	N
	ATOM	3726	CA	LEU	319	-0.814	-74.919	118.735	1.00 26.58	В	С
	ATOM	3727	СВ	LEU	319	0.302	-75.961	118.818	1.00 27.06	В	С
	ATOM	3728	CG	LEU	319	0.981	-76.399	117.524	1.00 28.14	В	С
25	MOTA	3729	CD1	LEU	319	1.450	-75.183	116.749	1.00 28.22	В	С
	ATOM	3730		LEU	319	2.156	-77.309	117.860	1.00 28.83	В	С
	ATOM	3731	C	LEU	319			118.036	1.00 26.66	В	C
	ATOM	3732	ō	LEU	319			116.815	1.00 25.59	В	ō
	ATOM	3733	N	ARG	320			118.826	1.00 26.81	В	N
30	ATOM	3734	CA	ARG	320			118.298	1.00 27.51	В	C
00	ATOM	3735	CB	ARG	320			119.388	1.00 27.31	В	c
	ATOM	3736	CG	ARG	320			118.892	1.00 30.21	В	c
	ATOM	3737	CD	ARG	320			117.808	1.00 34.14	В	C
		3738	NE		320			117.500	1.00 37.44	В	N
35	ATOM	3739	CZ	ARG ARG	320			118.118	1.00 41.13		
33	ATOM		-							В	C
	ATOM	3740		ARG	320			119.093 117.755	1.00 43.92	В	N
	ATOM	3741		ARG	320				1.00 43.56	В	N
	ATOM	3742	C	ARG	320			117.837	1.00 26.59	В	C
40	MOTA	3743	0	ARG	320			116.781	1.00 26.01	В	0
40	MOTA	3744	N	SER	321			118.635	1.00 25.46	В	N
	MOTA	3745	CA	SER	321			118.283	1.00 25.37	В	С
	MOTA	3746	CB	SER	321			119.356	1.00 25.01	В	C
	MOTA	3747	OG	SER	321			120.577	1.00 28.59	В	0
	MOTA	3748	С	SER	321			116.964	1.00 24.05	В	С
45	MOTA	3749	0	SER	321			116.097	1.00 23.58	В	0
	ATOM	3750	N	ILE	322			116.834	1.00 23.19	В	N
	MOTA	3751	CA	ILE	322			115.632	1.00 23.73	В	С
	MOTA	3752	CB	ILE	322			115.794	1.00 24.44	В	C
	ATOM	3753	CG2	! ILE	322			114.454	1.00 23.92	В	C
50	MOTA	3754	CG1	ILE	322			116.878	1.00 24.48	В	С
	ATOM	3755	CD1	LILE	322	0.068	-71.277	117.301	1.00 25.71	В	С
	ATOM	3756	С	ILE	322			114.418	1.00 23.91	В	C
	MOTA	3757	0	ILE	322			3 113.342	1.00 23.08	В	0
	ATOM	3758	N	ASN	323			114.601	1.00 24.23	В	N
55	MOTA	3759	CA	ASN	323			2 113.525	1.00 26.82	В	C
	MOTA	3760	СВ	ASN	323			114.033	1.00 29.48	В	č
	ATOM	3761	CG	ASN	323			112.929	1.00 23.40	В	C
	ATOM	3762		L ASN	323			113.058	1.00 32.74	В	0
	ALUM	3,02	JD.	154		2.420	, 0 . , 5		2.00 72.01	٠	9

	MOTA	3763	ND2		323		-77.208		1.00 34.14	В	N
	MOTA	3764	C	ASN	323		-74.764		1.00 27.38	В	С
	ATOM	3765	0	ASN	323		-74.546		1.00 26.41	В	0
_	MOTA	3766	N	GLU	324		-74.656		1.00 27.46	В	N
5	MOTA	3767	CA	GLU	324		-74.246		1.00 28.72	В	C
	MOTA	3768	CB	GLU	324	-8.867	-74.253	114.852	1.00 30.37	В	С
	MOTA	3769	CG	GLU	324	-8.979	-75.615	115.533	1.00 34.56	В	С
	ATOM	3770	CD	GLU	324	-9.872	-75.585	116.766	1.00 37.51	В	С
	MOTA	3771	OE1	GLU	324	-9.717	-74.660	117.598	1.00 38.60	В	0
10	ATOM	3772	OE2	GLU	324	-10.723	-76.492	116.909	1.00 39.78	В	0
	MOTA	3773	С	GLU	324		-72.846		1.00 27.55	В	C
	ATOM	3774	0	GLU	324		-72.574		1.00 28.12	В	Ō
	ATOM	3775	N	ALA	325		-71.959		1.00 25.61	В	N
	MOTA	3776	CA	ALA	325		-70.607		1.00 24.75	В	C
15	MOTA	3777	СВ	ALA	325		-69.763		1.00 25.33	В	Ċ
	MOTA	3778	C	ALA	325		-70.642		1.00 24.31	В	c
	ATOM	3779	Ö	ALA	325		-69.793		1.00 24.70	В	Ö
	ATOM	3780	N	TYR	326		-71.613		1.00 23.49	В	N
	ATOM	3781	CA	TYR	326		-71.763		1.00 23.45	В	C
20	ATOM	3782	СВ	TYR	326		-72.958		1.00 23.03	В	C
20	ATOM	3783	CG	TYR	326		-72.576		1.00 21.78	В	C
	ATOM	3784	CD1		326		-71.844		1.00 20.83	В	C
	MOTA	3785		TYR	326		-71.492		1.00 20.27		C
	MOTA	3786	CD2	TYR	326		-72.945		1.00 19.33	В	
25	ATOM	3787	CE2	TYR	326		-72.596			В	C
20	ATOM	3788	CZ	TYR	326		-71.872		1.00 19.56 1.00 19.09	В	С
	ATOM	3789	OH	TYR	326		-71.572			В	C
	MOTA	3790	C	TYR	326		-72.001		1.00 17.47 1.00 23.60	В	0
	ATOM	3791	0	TYR	326		-71.422			В	C
30	ATOM	3792	N	GLY	327		-72.862		1.00 23.00 1.00 23.42	В	0
00	ATOM	3793	CA	GLY	327		-73.174		1.00 23.42	В	И
	ATOM	3794	C	GLY	327		-71.936			В	C
	ATOM	3795	Ö	GLY	327		-71.773		1.00 26.46	В	C
		3796							1.00 25.91	В	0
35	ATOM ATOM	3797	N CA	TYR TYR	328		-71.056		1.00 27.71	В	N
33		3798	CB		328		-69.831		1.00 29.17	В	C
	ATOM		CG	TYR	328		-69.077		1.00 30.61	В	C
	ATOM	3799		TYR	328		-67.711		1.00 32.99	В	C
	MOTA	3800		TYR	328		-66.618		1.00 33.59	В	C
40	ATOM	3801		TYR	328		-65.381		1.00 33.86	В	C
40	ATOM	3802	CD2		328		-67.524		1.00 34.14	В	C
	ATOM	3803	CE2		328		-66.281		1.00 34.54	В	С
	ATOM	3804	CZ	TYR	328			109.758	1.00 35.00	В	
	ATOM	3805	ОН	TYR	328		-64.002		1.00 35.60	В	0
4 =	ATOM	3806	С	TYR	328		-68.936		1.00 29.11	В	С
45	ATOM	3807	0	TYR	328		-68.309		1.00 28.12	В	0
	ATOM	3808	N	GLN	329		-68.878		1.00 29.12	В	N
	ATOM	3809	CA	GLN	329		-68.058		1.00 29.08	В	С
	MOTA	3810	CB	GLN	329		-68.119		1.00 27.70	В	С
	MOTA	3811	CG	GLN	329		-67.517		1.00 26.09	В	C
50	ATOM	3812	CD	GLN	329		-66.032		1.00 25.68	В	С
	MOTA	3813		GLN	329		-65.578		1.00 26.08	В	0
	MOTA	3814		GLN	329		-65.267		1.00 24.01	В	N
	MOTA	3815	С	GLN	329		-68.488		1.00 30.18	В	С
_	MOTA	3816	0	GLN	329	-7.785	-67.649	104.932	1.00 29.17	В	0
55	MOTA	3817	N	ILE	330	-7.508	-69.792	105.561	1.00 31.28	В	N
	MOTA	3818	CA	ILE	330		-70.323		1.00 34.04	В	C
	MOTA	3819	СВ	ILE	330	-7.138	-71.795	104.140	1.00 35.02	В	C
	MOTA	3820	CG2	ILE	330			105.155	1.00 36.98	В	Ċ

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	MOTA	3821	CG1		330		-72.370		1.00 36.93	В	С
	MOTA	3822	CD1		330		-73.776		1.00 37.83	В	С
	MOTA	3823		ILE	330		-70.233		1.00 34.84	В	С
_	MOTA	3824		ILE	330	-	-70.200		1.00 33.41	В	0
5	ATOM	3825		GLN	331	-10.064			1.00 36.52	В	N
	MOTA	3826		GLN	331	-11.435			1.00 39.11	В	С
	MOTA	3827		GLN	331	-12.414			1.00 41.52	В	C
	MOTA	3828		GLN	331	-12.738			1.00 45.93	В	С
4.0	MOTA	3829		GLN	331	-11.691			1.00 48.62	В	С
10	MOTA	3830	OE1		331			106.400	1.00 50.00	В	0
	MOTA	3831	NE2		331			104.278	1.00 50.06	В	N
	MOTA	3832	С	GLN	331			103.741	1.00 38.77	В	С
	MOTA	3833	0	GLN	331			102.962	1.00 39.36	В	0
	MOTA	3834	N	HIS	332			104.408	1.00 38.40	В	N
15	MOTA	3835	CA	HIS	332			104.290	1.00 38.62	В	С
	MOTA	3836	CB	HIS	332			105.690	1.00 40.74	В	С
	MOTA	3837	CG	HIS	332			106.368	1.00 43.93	В	С
	MOTA	3838	CD2		332			107.075	1.00 44.81	В	С
	MOTA	3839	ND1		332			106.304	1.00 44.67	В	N
20	MOTA	3840	CE1		332			106.944	1.00 45.90	В	С
	MOTA	3841	NE2		332			107.419	1.00 45.60	В	N
	MOTA	3842	C	HIS	332			103.477	1.00 37.56	В	С
	MOTA	3843	0	HIS	332			103.146	1.00 36.45	В	0
	MOTA	3844	N	ILE	333			103.146	1.00 36.43	В	N
25	MOTA	3845	CA	ILE	333			102.369	1.00 35.55	В	С
	MOTA	3846	CB	ILE	333			103.194	1.00 35.40	В	C
	MOTA	3847	CG2	ILE	333			102.338	1.00 35.90	В	C
	ATOM	3848		ILE	333			104.392	1.00 35.38	В	C
	MOTA	3849		ILE	333			105.558	1.00 35.02	В	С
30	MOTA	3850	С	ILE	333			101.097	1.00 35.05	В	С
	MOTA	3851	0	ILE	333			101.108	1.00 34.80	В	0
	MOTA	3852	N	GLN	334			100.003	1.00 33.98	В	N
	MOTA	3853	CA	GLN	334		-66.072		1.00 33.55	В	С
0.5	MOTA	3854	CB	GLN	334		-65.284		1.00 34.76	В	C
35	MOTA	3855	CG	GLN	334		-65.984		1.00 37.74	В	C
	MOTA	3856	CD	GLN	334		-65.145		1.00 39.77	В	С
	MOTA	3857		GLN	334		-64.335		1.00 40.63	В	0
	MOTA	3858	NE2		334		-65.329		1.00 41.05	В	N
40	MOTA	3859	С	GLN	334		-66.101		1.00 31.83	В	C
40	MOTA	3860	0	GLN	334		-65.106		1.00 30.79	В	0
	ATOM	3861	N	GLY	335		-67.250		1.00 30.86	В	N
	ATOM	3862	CA	GLY	335		-67.356		1.00 30.05	В	
	MOTA	3863	C	GLY	335		-68.065		1.00 29.05	В	С
45	ATOM	3864	0	GLY	335		-68.703		1.00 28.90	В	0
45	ATOM	3865	N	LEU	336		-67.964		1.00 28.28	В	N
	MOTA	3866	CA	LEU	336			100.772	1.00 28.50	В	C
	MOTA	3867	СВ	LEU	336			102.189	1.00 26.88	В	C
	MOTA	3868	CG	LEU	336			. 102.775	1.00 26.76	В	C
50	MOTA	3869		LEU	336			104.198	1.00 25.90	В	C
50	MOTA	3870		LEU	336			102.781	1.00 26.43	В	C
	MOTA	3871	С	LEU	336			100.561	1.00 28.85	В	С
	MOTA	3872	0	LEU	336			100.761	1.00 28.06	В	0
	MOTA	3873	N	SER	337			100.149	1.00 29.32	В	N
	MOTA	3874	CA	SER	337		-72.208		1.00 30.16	В	С
55	MOTA	3875	CB	SER	337		-72.631		1.00 30.24	В	C
	ATOM	3876	OG	SER	337		-72.034			В	0
	ATOM	3877	С	SER	337		72.704		1.00 29.96	В	С
	MOTA	3878	0	SER	337	-3.374	-73.869	98.931	1.00 30.50	В	0

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	MOTA	3879	N	ALA	338		-71.828	97.996	1.00 29.75	В	N
	ATOM	3880	CA	ALA	338		-72.188	96.983	1.00 30.35	В	C
	ATOM	3881	СВ	ALA	338		-71.015	96.043	1.00 29.79	В	С
_	ATOM	3882	С	ALA	338		-72.615	97.615	1.00 30.54	В	С
5	ATOM	3883	0	ALA	338		-73.408	97.037	1.00 31.26	В	0
	MOTA	3884	N	MET	339	-0.736	-72.082	98.796	1.00 30.39	В	N
	MOTA	3885	CA	MET	339	0.499	-72.427	99.494	1.00 31.18	В	C
	MOTA	3886	CB	MET	339	0.906	-71.293	100.443	1.00 28.38	В	С
	ATOM	3887	CG	MET	339	1.428	-70.061	99.719	1.00 27.58	В	С
10	MOTA	3888	SD	MET	339	1.777	-68.662	100:804	1.00 25.78	В	S
	ATOM	3889	CE	MET	339		-67.932		1.00 26.72	В	C
	ATOM	3890	C	MET	339		-73.742		1.00 33.29	В	Č
	ATOM	3891	ō	MET	339		-74.253		1.00 32.44	В	ŏ
	MOTA	3892	N	MET	340		-74.283		1.00 36.51	В	N
15	ATOM	3893	CA	MET	340		-75.546		1.00 41.83	В	C
10	ATOM	3894	CB	MET	340		-75.740		1.00 41.83		C
	ATOM	3895	CG	MET	340		-76.879		1.00 45.44	B B	
	ATOM	3896	SD	MET	340		-76.795				C
									1.00 50.06	В	S
20	ATOM	3897	CE	MET	340		-78.542		1.00 49.89	В	C
20	ATOM	3898	С	MET	340		-76.639		1.00 44.42	В	С
	MOTA	3899	0	MET	340		-76.737	99.008	1.00 43.85	В	0
	ATOM	3900	Ŋ	PRO	341		-77.463		1.00 47.61	В	N
	ATOM	3901	CD	PRO	341		-77.444		1.00 48.17	В	C
0.5	MOTA	3902	CA	PRO	341		-78.593	99.902	1.00 50.66	В	С
25	MOTA	3903	CB	PRO	341		-79.506		1.00 49.85	В	С
	MOTA	3904	CG	PRO	341		-78.525	102.094	1.00 48.96	В	C
	MOTA	3905	С	PRO	341	0.470	-79.352	98.734	1.00 53.61	В	C
	ATOM	3906	0	PRO	341	1.190	-79.959	97.932	1.00 53.85	В	0
	MOTA	3907	N	LEU	342	-0.860	-79.329	98.640	1.00 56.59	В	N
30	MOTA	3908	CA	LEU	342	-1.595	-80.017	97.575	1.00 59.91	В	C
	MOTA	3909	CB	LEU	342	-0.835	-79.921	96.241	1.00 59.68	В	C
	ATOM	3910	CG	LEU	342	-1.238	-80.824	95.070	1.00 59.94	В	C
	ATOM	3911	CD1	LEU	342	-0.926	-80.135	93.752	1.00 59.43	В	C
	ATOM	3912	CD2	LEU	342	-0.506	-82.154	95.171	1.00 59.60	В	С
35	ATOM	3913	С	LEU	342	-1.845	-81.475	97.949	1.00 62.30	В	С
	ATOM	3914	0	LEU	342	-2.944	-81.997		1.00 62.71	В	0
	MOTA	3915	N	LEU	343	-0.821	-82.126		1.00 64.97	В	N
	ATOM	3916	CA	LEU	343		-83.516		1.00 67.52	В	C
	ATOM	3917	CB	LEU	343		-83.961		1.00 67.63	В	Č
40	ATOM	3918	CG	LEU	343		-83.566		1.00 67.80	В	č
. •	ATOM	3919		LEU	343		-84.029		1.00 67.88	В	Č
	ATOM	3920		LEU	343		-84.171		1.00 67.91	В	Č
	ATOM	3921	C	LEU	343			99.932	1.00 68.96	В	Ċ
	ATOM	3922	ŏ	LEU	343		-84.362		1.00 68.99	В	Ö
45	ATOM	3923	N	GLN	344			100.985	1.00 70.83	В	N
70	MOTA	3924	CA	GLN	344			102.048	1.00 70.63	В	C
	ATOM	3925	CB	GLN	344			102.048	1.00 72.82	В	
		3926	CG		344			103.128			С
	MOTA	3927	CD	GLN	344			104.428	1.00 72.55	В	C
50	MOTA			GLN					1.00 72.59	В	C
30	ATOM	3928		GLN	344			105.540	1.00 72.49	В	0
	MOTA	3929		GLN	344			103.832	1.00 72.52	В	N
	MOTA	3930	C	GLN	344			101.495	1.00 73.91	В	С
	MOTA	3931	0	GLN	344			101.740	1.00 74.16	В	0
e e	MOTA	3932	N	GLU	345			100.745	1.00 75.31	В	N
55	MOTA	3933	CA	GLU	345			100.166	1.00 76.49	В	C
	MOTA	3934	СВ	GLU	345			101.288	1.00 77.05	В	C
	ATOM	3935	CG	GLU	345			100.846	1.00 77.78	В	C
	MOTA	3936	CD	GLU	345	-8.705	-79.330	102.005	1.00 78.25	В	С

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	MOTA	3937	OE1		345	-8.867 -80.087 102.987 1.00 78.37 B	0
	MOTA	3938	OE2		345	-9.200 -78.184 101.937 1.00 78.57 B	0
	MOTA	3939	C	GLU	345	-5.258 -79.557 99.193 1.00 76.87 B	C
_	MOTA	3940	0	GLU	345	-5.484 -79.744 97.978 1.00 77.22 B	0
5	MOTA	3941	OXT		345	-4.802 -78.490 99.657 1.00 77.25 B	0
	TER	3942		GLU	345	В	
	MOTA	3943	CB	PRO	103	17.203 -24.177 122.780 1.00 92.75 C	С
	MOTA	3944	CG	PRO	103	15.916 -24.008 121.973 1.00 93.01 C	C
40	MOTA	3945	C	PRO	103	16.591 -26.001 124.396 1.00 92.31 C	С
10	MOTA	3946	0	PRO	103	15.433 -26.410 124.489 1.00 92.32 C	0
	MOTA	3947	N	PRO	103	16.430 -26.298 121.930 1.00 92.88 C	N
	MOTA	3948	CD	PRO	103	15.865 -25.248 121.064 1.00 93.06 C	С
	MOTA	3949	CA	PRO	103	17.200 -25.678 123.033 1.00 92.61 C	С
45	ATOM	3950	N	VAL	104	17.381 -25.812 125.450 1.00 91.79 C	N
15	ATOM	3951	CA	VAL	104	16.934 -26.091 126.812 1.00 91.07 C	С
	MOTA	3952	CB	VAL	104	17.809 -27.204 127.462 1.00 91.34 C	С
	MOTA	3953		VAL	104	17.551 -28.535 126.766 1.00 91.17 C	С
	MOTA	3954		VAL	104	19.289 -26.854 127.359 1.00 91.18 C	C
20	MOTA	3955	C	VAL	104	16.946 -24.835 127.691 1.00 90.40 C	С
20	MOTA	3956	0	VAL	104	17.957 -24.495 128.310 1.00 90.36 C	0
	MOTA	3957	N	GLN	105	15.803 -24.156 127.742 1.00 89.40 C	N
	MOTA	3958	CA	GLN	105	15.652 -22.928 128.519 1.00 88.14 C	С
	MOTA	3959	CB	GLN	105	14.585 -22.028 127.879 1.00 88.78 C	С
05	MOTA	3960	CG	GLN	105	15.024 -21.256 126.631 1.00 89.73 C	С
25	MOTA	3961	CD	GLN	105	15.344 -22.145 125.438 1.00 90.20 C	С
	MOTA	3962		GLN	105	16.377 -22.817 125.406 1.00 90.33 C	0
	MOTA	3963		GLN	105	14.454 -22.150 124.450 1.00 90.48 C	N
	MOTA	3964	C	GLN	105	15.274 -23.196 129.975 1.00 86.74 C	С
20	MOTA	3965	0	GLN	105	15.063 -24.345 130.372 1.00 87.04 C	0
30	ATOM	3966	N	LEU	106	15.193 -22.125 130.763 1.00 84.58 C	N
	MOTA	3967	CA	LEU	106	14.831 -22.211 132.176 1.00 82.19 C	C
	ATOM	3968	СВ	LEU	106	15.748 -23.199 132.907 1.00 82.41 C	С
	MOTA	3969	CG	LEU	106	15.400 -23.542 134.361 1.00 82.27 C	C
0.5	ATOM	3970		LEU	106	13.961 -24.027 134.458 1.00 82.05 C	С
35	MOTA	3971		LEU	106	16.356 -24.611 134.870 1.00 82.24 C	С
	MOTA	3972	C	LEU	106	14.920 -20.833 132.834 1.00 80.39 C	С
	ATOM	3973	0	LEU	106	15.976 -20.432 133.322 1.00 80.25 C	0
	ATOM	3974	N	SER	107	13.798 -20.118 132.833 1.00 78.12 C	N
40	ATOM	3975	CA	SER	107	13.698 -18.781 133.413 1.00 75.92 C	С
40	ATOM	3976	CB	SER	107	12.229 -18.437 133.670 1.00 75.50 C	С
	MOTA	3977	OG	SER	107	12.106 -17.218 134.377 1.00 74.70 C	0
	MOTA	3978	С	SER	107	14.484 -18.613 134.708 1.00 74.75 C	C
	MOTA	3979	0	SER	107	14.622 -19.552 135.491 1.00 74.59 C	0
45	MOTA	3980	N	LYS	108	14.998 -17.405 134.927 1.00 73.23 C	N
45	MOTA	3981	CA	LYS	108	15.761 -17.109 136.134 1.00 71.45 C	С
	MOTA	3982	CB	LYS	108	16.580 -15.825 135.954 1.00 71.64 C	C
	MOTA	3983	CG	LYS	108	15.752 -14.553 135.829 1.00 71.83 C	С
	MOTA	3984	CD	LYS	108	16.621 -13.317 135.598 1.00 72.24 C	C
	MOTA	3985	CE	LYS	108	17.070 -13.171 134.142 1.00 72.27 C	С
50	MOTA	3986	NZ	LYS	108	18.040 ~14.211 133.689 1.00 72.65 C	N
	MOTA	3987	С	LYS	108	14.828 -16.966 137.334 1.00 70.06 C	C
	MOTA	3988	0	LYS	108	15.241 -17.172 138.475 1.00 69.77 C	0
	MOTA	3989	N	GLU	109	13.572 -16.611 137.074 1.00 68.53 C	N
66	ATOM	3990	CA	GLU	109	12.589 -16.460 138.139 1.00 66.79 C	С
55	MOTA	3991	CB	GLU	109	11.342 -15.742 137.627 1.00 67.72 C	С
	MOTA	3992	CG	GLU	109	10.312 -15.496 138.709 1.00 69.22 C	С
	MOTA	3993	CD	GLU	109	9.117 -14.715 138.212 1.00 70.31 C	C
	MOTA	3994	OE1	GLU	109	9.308 -13.573 137.740 1.00 70.89 C	0

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	ATOM	3995	OE2		109			138.297	1.00 71.04	С	0
	MOTA	3996	С	GLU	109		-17.832		1.00 64.99	С	С
	MOTA	3997	0	GLU	109	-	-18.013		1.00 64.83	С	0
_	MOTA	3998	N	GLN	110		-18.799		1.00 62.56	C	N
5	MOTA	3999	CA	GLN	110		-20.138		1.00 60.51	С	C
	ATOM	4000	СВ	GLN	110		-20.952		1.00 60.12	С	С
	MOTA	4001	CG	GLN	110		-21.069		1.00 59.34	С	С
	MOTA	4002	CD	GLN	110		-21.510		1.00 58.88	С	C
4.0	MOTA	4003		GLN	110		-21.900		1.00 58.96	С	0
10	MOTA	4004		GLN	110		-21.438		1.00 58.28	С	N
	MOTA	4005	С	GLN	110		-20.818		1.00 59.45	С	С
	ATOM	4006	0	GLN	110			139.561	1.00 59.18	C	0
	MOTA	4007	N	GLU	111			138.538	1.00 57.98	С	N
	ATOM	4008	CA	GLU	111			139.139	1.00 56.65	C	С
15	MOTA	4009	CB	GLU	111			138.501	1.00 57.99	C	C
	MOTA	4010	CG	GLU	111			137.131	1.00 59.95	С	C
	MOTA	4011	CD	GLU	111		-20.842	136.885	1.00 61.39	С	С
	MOTA	4012		GLU	111			137.121	1.00 61.89	C	0
	MOTA	4013		GLU	111			136.453	1.00 61.84	С	0
20	MOTA	4014	С	GLU	111			140.613	1.00 54.63	С	C
	ATOM	4015	0	GLU	111			141.478	1.00 54.15	С	0
	MOTA	4016	N	GLU	112			140.881	1.00 52.67	С	N
	ATOM	4017	CA	GLU	112			142.240	1.00 51.06	С	С
	MOTA	4018	CB	GLU	112			142.209	1.00 52.04	C	С
25	MOTA	4019	CG	GLU	112			143.560	1.00 53.83	C	C
	MOTA	4020	CD	GLU	112			144.552	1.00 55.38	C	С
	MOTA	4021		GLU	112			144.575	1.00 56.05	С	0
	ATOM	4022	OE2	GLU	112	14.523	-15.527	145.324	1.00 56.06	С	0
	ATOM	4023	C	GLU	112			143.022	1.00 48.98	C	С
30	MOTA	4024	0	GLU	112	13.994	-20.042	144.149	1.00 48.54	C	0
	MOTA	4025	N	LEU	113			142.409	1.00 46.59	С	N
	MOTA	4026	CA	LEU	113	11.582	-20.852	143.013	1.00 43.90	C	C
	MOTA	4027	CB	LEU	113			142.030	1.00 43.46	С	C
	MOTA	4028	CG	LEU	113			142.419	1.00 42.93	C	С
35	MOTA	4029		LEU	113			143.685	1.00 41.97	С	С
	MOTA	4030		LEU	113			141.272	1.00 42.66	C	C
	MOTA	4031	С	LEU	113			143.396	1.00 42.33	C	C
	MOTA	4032	0	LEU	113			144.502	1.00 41.47	С	0
4.0	MOTA	4033	N	ILE	114			142.476	1.00 41.11	С	N
40	MOTA	4034	CA	ILE	114			142.766	1.00 40.20	C	С
	MOTA	4035	CB	ILE	114			141.521	1.00 39.72	С	C
	ATOM	4036		ILE	114			141.902	1.00 39.64	С	
	MOTA	4037		ILE	114			140.458	1.00 40.14	С	С
4.5	MOTA	4038		ILE	114			139.226	1.00 39.70	С	C
45	ATOM	4039	С	ILE	114			143.914	1.00 39.98	С	С
	ATOM	4040	0	ILE	114			144.782	1.00 39.82	С	0
	MOTA	4041	N	ARG	115			143.916	1.00 39.67	С	N
	MOTA	4042	CA	ARG	115			144.956	1.00 39.67	С	C
	MOTA	4043	CB	ARG	115			144.670	1.00 41.84	С	С
50	MOTA	4044	CG	ARG	115			145.596	1.00 45.10	С	С
	MOTA	4045	CD	ARG				146.953	1.00 47.84	C	C
	MOTA	4046	NE	ARG				146.880	1.00 50.53	С	N
	MOTA	4047	CZ	ARG				147.901	1.00 52.18	С	С
	MOTA	4048		. ARG				149.086	1.00 52.65	С	N
55	MOTA	4049		ARG				147.739		C	N
	MOTA	4050	С	ARG				146.315		С	С
	MOTA	4051	0	ARG				147.260		С	0
	MOTA	4052	N	THR	116	14.552	-21.639	146.401	1.00 36.28	С	N

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	ATOM	4053	CA	THR	116		-21.512		1.00 34.60	С	C
	MOTA	4054	CB	THR	116		-20.439		1.00 35.71	C	С
	MOTA	4055	OG1		116		-20.351		1.00 37.91	C	0
_	MOTA	4056		THR	116		-20.776		1.00 37.42	C	С
5	MOTA	4057	C	THR	116		-22.864		1.00 32.61	C	С
	MOTA	4058	0	THR	116	13.281	-23.189	149.287	1.00 31.75	C	0
	ATOM	4059	N	LEU	117	12.692	-23.660	147.171	1.00 30.08	C	N
	MOTA	4060	CA	LEU	117	12.138	-24.971	147.518	1.00 28.36	C	Ç
	MOTA	4061	CB	LEU	117	11.458	-25.614	146.302	1.00 26.76	С	С
10	ATOM	4062	CG	LEU	117	10.176	-24.967	145.769	1.00 25.75	C	С
	MOTA	4063	CD1	LEU	117	9.678	-25.739	144.568	1.00 25.12	C	С
	ATOM	4064	CD2	LEU	117		-24.940		1.00 24.24	C	С
	MOTA	4065	С	LEU	117		-25.904		1.00 28.17	С	С
	MOTA	4066	Ō	LEU	117		-26.608		1.00 27.39	Ċ	0
15	ATOM	4067	N	LEU	118		-25.908		1.00 27.82	C	N
. •	ATOM	4068	CA	LEU	118		-26.751		1.00 27.94	Č	C
	ATOM	4069	СВ	LEU	118		-26.639		1.00 28.74	č	Č
	ATOM	4070	CG	LEU	118			145.661	1.00 30.12	Ċ	Ċ
	ATOM	4071		LEU	118			145.087	1.00 30.71	c	C
20	ATOM	4072		LEU	118			144.556	1.00 30.71	C	c
20	MOTA	4073	C	LEU	118			149.105	1.00 31.14	c	C
		4074		LEU	118			149.103		C	Ö
	MOTA		0		119				1.00 26.71		
	MOTA	4075	N	GLY				149.366	1.00 26.49	C	N
25	ATOM	4076	CA	GLY	119			150.655	1.00 26.44	C	C
25	ATOM	4077	C	GLY	119			151.782	1.00 26.03	C	C
	ATOM	4078	0	GLY	119			152.714	1.00 25.94	C	0
	ATOM	4079	N	ALA	120			151.694	1.00 25.49	C	N
	ATOM	4080	CA	ALA	120			152.707	1.00 24.69	С	C
20	ATOM	4081	СВ	ALA	120			152.329	1.00 24.54	C	С
30	MOTA	4082	C	ALA	120			152.858	1.00 24.27	C	С
	ATOM	4083	0	ALA	120			153.972	1.00 23.29	C	0
	MOTA	4084	N	HIS	121			151.730	1.00 23.87	C	N
	MOTA	4085	CA	HIS	121			151.733	1.00 23.41	C	C
o.e.	MOTA	4086	CB	HIS	121			150.302	1.00 23.48	C	C
35	MOTA	4087	CG	HIS	121			150.197	1.00 23.06	C	C
	MOTA	4088		HIS	121			150.383	1.00 22.98	C	С
	ATOM	4089		HIS	121			149.890	1.00 24.73	С	N
	MOTA	4090		HIS	121			149.890	1.00 24.22	С	С
4.0	MOTA	4091		HIS	121			150.187	1.00 24.19	C	N
40	MOTA	4092	С	HIS	121			152.394	1.00 23.45	C	C
	ATOM	4093	0	HIS	121			153.230	1.00 22.84	С	0
	MOTA	4094	N	THR	122			152.018	1.00 23.17	С	
	MOTA	4095	CA	THR	122			152.591	1.00 23.53	С	C
	MOTA	4096	CB	THR	122	18.379	-28.847	151.918	1.00 23.68	С	С
45	ATOM	4097	OG1	THR	122	18.483	-29.298	150.560	1.00 23.80	С	0
	MOTA	4098	CG2	THR	122			152.657	1.00 23.99	C	C
	MOTA	4099	С	THR	122	17.193	-29.309	154.106	1.00 23.12	С	С
	MOTA	4100	0	THR	122	17.526	-30.215	154.872	1.00 22.20	С	0
	MOTA	4101	N	ARG	123	16.826	-28.110	154.535	1.00 22.95	C	N
50	MOTA	4102	CA	ARG	123	16.843	-27.780	155.954	1.00 23.76	С	C
	MOTA	4103	СВ	ARG	123	16.591	-26.275	156.131	1.00 25.69	С	C
	ATOM	4104	CG	ARG	123			157.549	1.00 28.57	С	C
	ATOM	4105	CD	ARG	123			157.692	1.00 30.95	c	č
	ATOM	4106	NE	ARG	123			156.775	1.00 33.62	č	
55	ATOM	4107	CZ	ARG	123	18.190		156.996	1.00 34.51	c	C
	ATOM	4108		L ARG	123			158.111	1.00 33.79	C	N
	ATOM	4109		2 ARG	123			156.095	1.00 33.79	C	
	ATOM	4110	C	ARG	123			156.819	1.00 34.90	C	
	27 OF		_			040	-20.300	, 10.013	1.00 22.09	C	C

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		4111	_		100		00 045				_
	ATOM	4111	0	ARG	123		-28.945		1.00 22.11	C	0
	MOTA	4112	N	HIS	124		-28.824		1.00 21.71	С	N
	MOTA	4113	CA	HIS	124		-29.483		1.00 21.10	C	C
_	MOTA	4114	CB	HIS	124		-28.565		1.00 20.20	C	С
5	MOTA	4115	CG	HIS	124		-27.207		1.00 19.38	C	С
	MOTA	4116	CD2	HIS	124	13.214	-26.818	158.909	1.00 17.97	C	С
	MOTA	4117	ND1	HIS	124	12.473	-26.050	157.002	1.00 19.43	С	N
	ATOM	4118	CE1	HIS	124	12.832	-25.008	157.731	1.00 18.44	С	С
	MOTA	4119	NE2	HIS	124	13.285	-25.447	158.892	1.00 18.55	С	N
10	ATOM	4120	С	HIS	124	13.108	-30.888	156.790	1.00 21.31	C	C
	ATOM	4121	0	HIS	124		-31.555		1.00 20.78	Ċ	Ō
	MOTA	4122	N	MET	125		-31.352		1.00 21.02	Č	N
	ATOM	4123	CA	MET	125		-32.665		1.00 21.61	Č	C
	ATOM	4124	СВ	MET	125		-32.478		1.00 22.63	C	C
15	MOTA	4125	CG	MET	125		-31.555		1.00 24.18	C	C
	ATOM	4126	SD	MET	125		-31.737		1.00 24.18		s
	ATOM	4127	CE	MET	125		-31.067			C	
	MOTA	4128	C	MET	125		-33.673		1.00 26.67	C	C
	ATOM	4129							1.00 20.49	C	C
20			0	MET	125		-34.860		1.00 19.83	С	0
20	ATOM	4130	N	GLY	126		-33.195		1.00 20.13	С	N
	MOTA	4131	CA	GLY	126		-34.063		1.00 19.93	С	С
	ATOM	4132	C	GLY	126		-35.193		1.00 20.19	С	С
	MOTA	4133	0	GLY	126		-36.333		1.00 19.39	C	0
0.5	MOTA	4134	N	THR	127		-34.893		1.00 20.00	С	N
25	MOTA	4135	CA	THR	127		-35.929		1.00 20.76	С	С
	MOTA	4136	CB	THR	127	18.575	-35.664	156.155	1.00 20.73	C	С
	MOTA	4137	OG1	THR	127	18.755	-34.329	156.638	1.00 20.90	С	0
	MOTA	4138	CG2	THR	127	19.349	-35.824	154.850	1.00 19.98	С	С
	MOTA	4139	С	THR	127	16.369	-36.086	157.267	1.00 20.55	С	C
30	MOTA	4140	0	THR	127		-36.534		1.00 19.57	С	0
	ATOM	4141	N	MET	128		-35.739		1.00 20.36	c	N
	ATOM	4142	CA	MET	128		-35.846		1.00 20.82	Č	C
	ATOM	4143	CB	MET	128		-35.193		1.00 21.34	C	Č
	ATOM	4144	CG	MET	128		-35.955		1.00 21.05	c	Ċ
35	ATOM	4145	SD	MET	128			157.462	1.00 20.70	C	s
•	ATOM	4146	CE	MET	128		-36.229		1.00 20.70	c	C
	ATOM	4147	C	MET	128			159.019	1.00 21.09	C	
	ATOM	4148	ō	MET	128			160.207	-		C
	ATOM	4149	N	PHE	129		-38.232		1.00 19.74	C	0
40	ATOM	4150	CA	PHE	129				1.00 21.12	C	N
70		4151						158.386	1.00 21.43	C	С
	MOTA		CB	PHE	129			157.089	1.00 23.16	С	C
	ATOM	4152	CG	PHE	129			156.385	1.00 25.47	С	_
	ATOM	4153		PHE	129		-41.413		1.00 25.73	С	С
AE	ATOM	4154		PHE	129			155.298	1.00 26.68	С	C
45	ATOM	4155		PHE	129	17.637	-41.465	156.200	1.00 28.33	С	С
	MOTA	4156		PHE	129			154.661	1.00 27.60	С	C
	MOTA	4157	CZ	PHE	129			155.112	1.00 28.66	С	C
	ATOM	4158	С	PHE	129	15.229	-40.150	159.300	1.00 20.85	С	C
	MOTA	4159	0	PHE	129			159.983	1.00 19.76	С	0
50	MOTA	4160	N	GLU	130			159.314	1.00 20.93	С	N
	ATOM	4161	CA	GLU	130	17.490	-39.860	160.153	1.00 22.74	С	С
	ATOM	4162	CB	GLU	130			159.875	1.00 24.78	C	Ċ
	MOTA	4163	CG	GLU	130			158.446	1.00 27.92	Č	Č
	ATOM	4164	CD	GLU	130			158.231	1.00 29.79	č	C
55	MOTA	4165		GLU	130			159.051	1.00 30.38	c	o
	ATOM	4166		GLU	130			157.236	1.00 30.83	c	0
	ATOM	4167	C	GLU				161.637	1.00 30.83	C	C
	MOTA	4168	Ö	GLU				162.445	1.00 21.42	C	
	*** OF	*100	•	300	# J U	-,.000	/	102.443	1.00 21.10	U	0

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	MOTA	4169	N	GLN	131		-39.001		1.00 20.13	С	N
	MOTA	4170	CA	GLN	131	15.770	-38.843	163.382	1.00 20.44	C	С
	MOTA	4171	CB	GLN	131		-37.442		1.00 20.12	C	С
_	MOTA	4172		GLN	131		-36.305		1.00 22.04	С	С
5	ATOM	4173	CD	GLN	131		-36.478		1.00 22.44	C	С
	MOTA	4174	OE1	GLN	131	17.630	-36.726	165.156	1.00 20.89	С	0
	ATOM	4175	NE2	GLN	131	18.588	-36.347	163.149	1.00 21.74	С	N
	ATOM	4176	С	GLN	131	14.766	-39.880	163.878	1.00 20.14	C	С
	MOTA	4177	0	GLN	131	14.500	-39.947	165.078	1.00 19.21	С	0
10	MOTA	4178	N	PHE	132	14.211	-40.686	162.970	1.00 18.92	С	N
	ATOM	4179	CA	PHE	132	13.235	-41.709	163.360	1.00 18.44	С	С
	MOTA	4180	CB	PHE	132	12.850	-42.587	162.162	1.00 17.81	С	С
	MOTA	4181	CG	PHE	132	12.019	-41.883	161.104	1.00 18.33	С	С
	ATOM	4182	CD1	PHE	132	11.560	-40.576	161.286	1.00 18.03	С	С
15	MOTA	4183	CD2	PHE	132	11.692	-42.547	159.920	1.00 17.90	С	С
	ATOM	4184	CE1	PHE	132		-39.944		1.00 18.71	С	С
	ATOM	4185	CE2	PHE	132	10.927	-41.932	158.936	1.00 18.59	С	С
	ATOM	4186	CZ	PHE	132		-40.624		1.00 18.77	С	С
	MOTA	4187	С	PHE	132		-42.607		1.00 18.64	C	C
20	MOTA	4188	0	PHE	132		-43.061		1.00 16.29	Ċ	Ō
	ATOM	4189	N	VAL	133	15.081	-42.866	164.446	1.00 18.83	Č	N
	ATOM	4190	CA	VAL	133	15.699	-43.714	165.450	1.00 19.85	Č	Ċ
	ATOM	4191	CB	VAL	133		-43.995		1.00 20.35	Ċ	Ċ
	ATOM	4192	CG1	VAL	133		-42.709		1.00 19.58	c	Ċ
25	ATOM	4193	CG2	VAL	133			166.034	1.00 20.81	Č	Ċ
	MOTA	4194	С	VAL	133		-43.124		1.00 20.00	č	Č
	ATOM	4195	0	VAL	133			167.841	1.00 20.01	Ċ	ō
	ATOM	4196	N	GLN	134			166.957	1.00 20.18	Č	N
	MOTA	4197	CA	GLN	134			168.261	1.00 19.68	Č	c
30	ATOM	4198	СВ	GLN	134			168.187	1.00 20.98	Č	Č
	ATOM	4199	CG	GLN	134			167.598	1.00 21.75	č	č
	ATOM	4200	CD	GLN	134			168.411	1.00 24.09	č	č
	ATOM	4201	OE1	GLN	134			167.982	1.00 25.24	Ċ	ō
	ATOM	4202	NE2		134			169.577	1.00 23.26	c	И
35	ATOM	4203	С	GLN	134			168.847	1.00 19.44	Č	C
	ATOM	4204	0	GLN	134			169.909	1.00 18.53	č	ŏ
	ATOM	4205	N	PHE	135			168.178	1.00 17.76	č	N
	MOTA	4206	CA	PHE	135			168.656	1.00 17.30	Č	c
	ATOM	4207	СВ	PHE	135			167.599	1.00 16.60	č	C
40	ATOM	4208	CG	PHE	135			167.467	1.00 15.94	Č	Č
	ATOM	4209	CD1		135			168.406	1.00 15.05	c	c
	MOTA	4210	CD2	PHE	135			166.432	1.00 16.21	C	
	ATOM	4211		PHE	135			168.323	1.00 16.31	č	Ċ
	ATOM	4212		PHE	135			166.337	1.00 15.93	C	Ċ
45	ATOM	4213	CZ	PHE	135			167.288	1.00 15.48	C	c
	ATOM	4214	C	PHE	135			169.072	1.00 17.74	č	C
	ATOM	4215	ō	PHE	135			168.587	1.00 16.28	c	Ö
	ATOM	4216	N	ARG	136			169.982	1.00 18.46	č	N
	ATOM	4217	CA	ARG	136			170.522	1.00 20.06	č	C
50	ATOM	4218	СВ	ARG	136			171.516	1.00 20.96	Č	Č
	ATOM	4219	CG	ARG	136			172.580	1.00 24.90	č	C
	ATOM	4220	CD	ARG	136			173.505	1.00 26.50	C	C
	ATOM	4221	NE	ARG	136			172.903	1.00 27.43	c	N
	ATOM	4222	CZ	ARG	136			173.598	1.00 27.43	c	C
55	ATOM	4223		ARG	136			174.903	1.00 27.65	c	N
- •	ATOM	4224		ARG	136			173.001	1.00 27.78	C	
	ATOM	4225	C	ARG	136			169.489	1.00 27.78	C	C N
	ATOM	4226	Ö	ARG	136			169.585	1.00 19.43	C	0
	ALON		9	MA	200	10.042	-31.000	109.505	1.00 20.16	C	U

	ATOM	4227	N	PRO	137		-46.614		1.00 18.91	С	N
	MOTA	4228	CD	PRO	137		-46.014		1.00 18.50	С	С
	MOTA	4229	CA	PRO	137		-47.637		1.00 18.39	С	С
_	MOTA	4230	CB	PRO	137				1.00 18.19	С	С
5	MOTA	4231	CG	PRO	137	13.844	-46.991	167.190	1.00 18.43	С	C
	MOTA	4232	С	PRO	137	11.677	-49.037	168.119	1.00 18.32	С	С
	ATOM	4233	0	PRO	137	12.612	-49.338	168.849	1.00 17.61	С	0
	ATOM	4234	N	PRO	138	10.690	-49.903	167.845	1.00 18.18	C	N
	ATOM	4235	CD	PRO	138		-49.665	167.165	1.00 18.09	Ċ	C
10	ATOM	4236	CA	PRO	138		-51.259		1.00 18.53	c	Ċ
	ATOM	4237	СВ	PRO	138		-51.905		1.00 19.26	Č	č
	ATOM	4238	CG	PRO	138		-50.729		1.00 20.92	C	C
	ATOM	4239	c	PRO	138			167.851	1.00 18.85	C	C
	ATOM	4240	Õ	PRO	138		-51.588		1.00 18.83	C	0
15	ATOM	4241	N	ALA	139		-52.914				
10	ATOM	4242	CA	ALA	139		-53.608		1.00 18.83	C	N
		4242	CB	ALA				168.170	1.00 19.23	C	C
	MOTA				139		-54.610	169.249	1.00 19.24	C	С
	ATOM	4244	C	ALA	139		-54.314		1.00 20.00	С	С
20	MOTA	4245	0	ALA	139		-54.355	166.090	1.00 19.18	С	0
20	MOTA	4246	N	HIS	140		-54.870	166.448	1.00 19.64	C	N
	MOTA	4247	CA	HIS	140		-55.586	165.180	1.00 20.90	С	С
	MOTA	4248	СВ	HIS	140		-56.340	165.073	1.00 19.67	C	С
	MOTA	4249	CG	HIS	140		-55.479	164.690	1.00 20.46	С	С
	MOTA	4250	CD2	HIS	140	9.320	-55.333	163.508	1.00 20.20	С	С
25	MOTA	4251	ND1	HIS	140	9.340	-54.632	165.577	1.00 19.33	C	N
	MOTA	4252	CE1	HIS	140	8.357	-54.000	164.959	1.00 20.25	C	С
	MOTA	4253	NE2	HIS	140	8.324	-54.406	163.704	1.00 20.65	С	N
	MOTA	4254	С	HIS	140	12.712	-54.727	163.927	1.00 21.31	C	С
	MOTA	4255	0	HIS	140		-55.259	162.827	1.00 21.71	Ċ	ō
30	ATOM	4256	N	LEU	141		-53.410	164.092	1.00 21.64	Ċ	N
	MOTA	4257	CA	LEU	141		-52.505		1.00 22.55	Č	C
	ATOM	4258	СВ	LEU	141		-51.156		1.00 21.20	č	č
	ATOM	4259	CG	LEU	141	10.852	-51.128		1.00 21.22	C	Ċ
	ATOM	4260		LEU	141		-49.687		1.00 19.35	C	Ċ
35	ATOM	4261		LEU	141		-51.742	162.274	1.00 20.05	C	C
00	ATOM	4262	C	LEU	141		-52.244				
	ATOM	4263	0	LEU	141	14.845			1.00 23.75	C	С
			-						1.00 23.76	C	0
	MOTA	4264	N	PHE	142		-52.677		1.00 24.92	C	N
40	ATOM	4265	CA	PHE	142		-52.413	163.377	1.00 27.83	С	С
40	ATOM	4266	СВ	PHE	142		-52.409	164.733	1.00 26.48	C	С
	MOTA	4267	CG	PHE	142			165.535	1.00 26.04	С	C
	ATOM	4268		PHE	142			166.508	1.00 26.71	С	С
	MOTA	4269		PHE	142			165.340	1.00 25.71	С	C
4 =	MOTA	4270		PHE	142			167.283	1.00 26.31	С	C
45	MOTA	4271		PHE	142			166.107	1.00 25.93	С	C
	MOTA	4272	CZ	PHE	142			167.080	1.00 24.84	C	С
	MOTA	4273	С	PHE	142			162.428	1.00 29.94	С	С
	ATOM	4274	0	PHE	142			162.376	1.00 29.88	С	0
	MOTA	4275	N	ILE	143	18.547	-52.700	161.704	1.00 32.51	С	N
50	ATOM	4276	CA	ILE	143	19.461	-53.355	160.756	1.00 35.12	С	С
	MOTA	4277	CB	ILE	143			161.497	1.00 35.28	C	Ċ
	ATOM	4278		ILE	143			160.519	1.00 34.76	c	Č
	ATOM	4279		ILE	143			162.632	1.00 35.53	c	c
	ATOM	4280		ILE	143			162.192	1.00 36.82	C	c
55	ATOM	4281	C	ILE	143			159.657	1.00 36.64	C	c
	ATOM	4282	ŏ	ILE	143			158.483	1.00 36.19	C	0
	MOTA	4283	N	HIS	144			160.030	1.00 38.19	C	
	ATOM	4284	CA	HIS	144			159.082			N
	AION	4404	CA		722	11.423	-30.08/	133.002	1.00 39.85	С	С

	3 0001	4005	<b>a</b> n				67 430				_
	ATOM	4285	CB	HIS	144		-57.439		1.00 42.16	С	С
	MOTA	4286	CG	HIS	144		-58.290		1.00 44.99	C	C
	MOTA	4287	CD2		144		-58.062		1.00 45.85	C	C
_	MOTA	4288	ND1		144		-59.550		1.00 46.31	С	N
5	MOTA	4289	CE1		144		-60.061		1.00 46.80	C	C
	MOTA	4290	NE2	HIS	144	16.008	-59.178	156.344	1.00 46.60	С	N
	MOTA	4291	С	HIS	144	16.073	-56.302	159.728	1.00 39.44	C	С
	MOTA	4292	0	HIS	144	15.977	-56.886	160.809	1.00 38.57	С	0
	MOTA	4293	N	HIS	145	15.026	-55.812	159.075	1.00 39.00	С	N
10	MOTA	4294	CA	HIS	145	13.693	-55.972	159.615	1.00 39.08	С	C
	MOTA	4295	CB	HIS	145		-55.528		1.00 38.43	Č	Ċ
	ATOM	4296	CG	HIS	145		-55.408		1.00 38.37	Ċ	Č
	ATOM	4297	CD2	HIS	145		-54.322		1.00 37.79	Ċ	Ċ
	MOTA	4298	ND1		145		-56.501		1.00 37.09	Č	N
15	ATOM	4299		HIS	145		-56.095		1.00 37.03	C	C
. •	ATOM	4300		HIS	145		-54.777		1.00 37.12	C	N
	ATOM	4301	C	HIS	145		-57.436		1.00 37.13		C
	ATOM	4302	Ö	HIS	145		-58.343			С	
	MOTA	4302	N	GLN					1.00 38.89	С	0
20					146		-57.650		1.00 38.87	C	N
20	MOTA	4304	CA	GLN	146		-58.984		1.00 38.47	C	C
	ATOM	4305	CB	GLN	146		-59.213		1.00 39.68	С	С
	ATOM	4306	CG	GLN	146		-59.681		1.00 41.08	C	С
	MOTA	4307	CD	GLN	146		-59.921		1.00 43.84	C	С
05	MOTA	4308	OE1		146		-58.976		1.00 45.31	С	0
25	MOTA	4309	NE2		146		-61.188		1.00 44.12	C	N
	ATOM	4310	С	GLN	146		-59.107		1.00 37.20	С	С
	MOTA	4311	0	GLN	146		-58.161		1.00 37.37	С	0
	MOTA	4312	N	PRO	147		-60.284		1.00 35.99	C	N
	MOTA	4313	CD	PRO	147	11.236	-61.547	161.151	1.00 35.90	C	С
30	ATOM	4314	CA	PRO	147	9.184	-60.423	161.788	1.00 34.04	С	С
	MOTA	4315	CB	PRO	147		-61.840		1.00 34.54	Ċ	C
	ATOM	4316	CG	PRO	147		-62.558		1.00 35.92	Ċ	Ċ
	ATOM	4317	С	PRO	147		-60.213		1.00 32.25	Č	Č
	ATOM	4318	0	PRO	147		-60.741		1.00 31.73	Č	ŏ
35	ATOM	4319	N	LEU	148		-59.406		1.00 29.88	Ċ	N
	ATOM	4320	CA	LEU	148		-59.167		1.00 26.83	C	C
	ATOM	4321	СВ	LEU	148		-58.001		1.00 27.33	C	c
	ATOM	4322	CG	LEU	148		-57.566		1.00 28.42	C	C
	ATOM	4323		LEU	148		-57.134		1.00 28.52	C	C
40	ATOM	4324	CD2		148		-56.420		1.00 28.07		2
.0	ATOM	4325	C	LEU	148		-60.478		1.00 24.35	C	C
	ATOM	4326	0	LEU	148					C	C
	ATOM	4327	N	PRO	149	5.809	-61.013 -61.037	164.344	1.00 22.45	C	0
			CD						1.00 22.72	C	N
45	ATOM	4328		PRO	149		-60.668		1.00 21.82	C	C
40	ATOM	4329	CA	PRO	149		-62.301		1.00 21.92	С	С
	ATOM	4330	CB	PRO	149		-62.579		1.00 20.94	С	С
	ATOM	4331	CG	PRO	149		-61.986		1.00 22.25	С	C
	MOTA	4332	С	PRO	149		-62.222		1.00 21.18	С	С
	MOTA	4333	0	PRO	149		-61.164		1.00 21.31	С	0
50	ATOM	4334	N	THR	150		-63.362		1.00 20.81	C	N
	ATOM	4335	CA	THR	150		-63.470		1.00 20.80	C	C
	ATOM	4336	CB	THR	150	2.152	-64.950	166.233	1.00 19.51	С	C
	MOTA	4337	OG1	THR	150	2.553	-65.453	164.951	1.00 17.95	С	0
	MOTA	4338	CG2	THR	150			166.404	1.00 19.20	С	Ċ
55	MOTA	4339	С	THR	150			167.532	1.00 21.88	Č	c
	MOTA	4340	0	THR	150			167.336	1.00 21.66	Č	ŏ
	MOTA	4341	N	LEU	151			168.751	1.00 21.61	c	N
	ATOM	4342	CA	LEU	151		-62.669		1.00 21.32	C	C
							02.009	107.751	x.00 2x.32	_	_

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	MOTA	4343	СВ	LEU	151	1.369	-63.796	170.964	1.00 20.79	С	С
	ATOM	4344	CG	LEU	151		-64.995		1.00 20.54	Ċ	C
	MOTA	4345	CD1		151		-66.025		1.00 20.72	C	C
	ATOM	4346	CD2	LEU	151		-64.530		1.00 19.30	Ċ	Ċ
5	MOTA	4347	С	LEU	151		-61.455		1.00 21.12	Ċ	Ċ
	ATOM	4348	0	LEU	151		-61.012		1.00 21.25	č	ō
	ATOM	4349	N	ALA	152		-60.916		1.00 19.97	Č	N
	ATOM	4350	CA	ALA	152		-59.751		1.00 19.76	Č	C
	MOTA	4351	СВ	ALA	152		-59.516	-	1.00 18.98	c	Č
10	ATOM	4352	С	ALA	152		-58.506		1.00 19.64	Č	Č
	ATOM	4353	ō	ALA	152		-58.267		1.00 18.40	c	ŏ
	ATOM	4354	N	PRO	153		-57.705		1.00 19.90	c	N
	ATOM	4355	CD	PRO	153			172.754	1.00 20.17	c	c
	ATOM	4356	CA	PRO	153		-56.495		1.00 20.12	C	c
15	ATOM	4357	CB	PRO	153		-55.865		1.00 20.62	c	c
	ATOM	4358	CG	PRO	153		-57.067		1.00 20.46	č	Č
	ATOM	4359	c	PRO	153			170.286	1.00 20.10	Č	Č
	ATOM	4360	Ö	PRO	153			169.989	1.00 19.34	C	Ö
	ATOM	4361	N	VAL	154			169.626	1.00 21.19	Č	N
20	ATOM	4362	CA	VAL	154			168.522	1.00 22.10	c	C
	ATOM	4363	CB	VAL	154			167.388	1.00 23.65	C	Ċ
	MOTA	4364		VAL	154			167.869	1.00 24.74	C	C
	ATOM	4365		VAL	154			166.172	1.00 24.74	C	C
	ATOM	4366	C	VAL	154			169.026	1.00 21.26	C	C
25	ATOM	4367	Ö	VAL	154			168.296	1.00 21.70	C	ō
	ATOM	4368	N	LEU	155			170.284	1.00 20.70	C	Ŋ
	MOTA	4369	CA	LEU	155			170.902	1.00 20.70	C	C
	ATOM	4370	СВ	LEU	155			172.393	1.00 19.93	C	C
	ATOM	4371	CG	LEU	155			173.250	1.00 21.72	c	c
30	ATOM	4372		LEU	155			172.660	1.00 20.06	C	C
•	ATOM	4373		LEU	155			174.681	1.00 20.00	C	c
	ATOM	4374	C	LEU	155			170.726	1.00 18.83	c	c
	ATOM	4375	ŏ	LEU	155			170.406	1.00 10.03	C	ŏ
	ATOM	4376	N	PRO	156			170.942	1.00 13.02	C	N
35	ATOM	4377	CD	PRO	156			171.571	1.00 17.61	C	C
-	MOTA	4378	CA	PRO	156			170.754	1.00 17.01	c	č
	ATOM	4379	СВ	PRO	156			171.141	1.00 17.22	C	c
	ATOM	4380	CG	PRO	156			172.205	1.00 10.70	C	C
	ATOM	4381	C	PRO	156			169.314	1.00 17.71	Č	C
40	ATOM	4382	Ö	PRO	156			169.093	1.00 16.22	c	ŏ
	ATOM	4383	N	LEU	157			168.342	1.00 16.66	c	N
	ATOM	4384	CA	LEU	157			166.942	1.00 17.15	Č	Ĉ
	ATOM	4385	СВ	LEU	157			166.019	1.00 15.68	C	Č
	MOTA	4386	CG	LEU	157			164.516	1.00 14.50	C	Č
45	ATOM	4387		LEU	157			164.153	1.00 12.24	C	Č
. •	ATOM	4388		LEU	157			163.714	1.00 11.45	C	Č
	ATOM	4389	C	LEU	157			166.688	1.00 16.47	c	c
	ATOM	4390	ŏ	LEU	157			166.016	1.00 16.30	C	ō
	ATOM	4391	N	VAL	158			167.229	1.00 16.36	Ċ	N
50	ATOM	4392	CA	VAL	158			167.088	1.00 16.91	c	C
•	ATOM	4393	СВ	VAL	158			167.762	1.00 17.14	C	c
	ATOM	4394		VAL	158			167.732	1.00 15.19	c	C
	ATOM	4395		VAL	158			167.046	1.00 15.59	Č	C
	ATOM	4396	C	VAL	158			167.727	1.00 13.33	c	c
55	ATOM	4397	Ö	VAL	158			167.154	1.00 17.43	C	0
	MOTA	4398	N	THR	159			168.919	1.00 17.33	C	N
	MOTA	4399	CA	THR	159			169.605	1.00 13.70	C	C
	MOTA	4400	СВ	THR	159			170.976	1.00 17.03	C	C
	-11-01-1	0 0			100	3.043		1.0.3/0	1.00 10.94	C	C

		4401	001	mun	150	3 040	46 505	171 776	1 00 15 06	_	_
	MOTA	4401	0G1		159		-46.585		1.00 15.26	С	0
	ATOM	4402		THR	159		-44.700		1.00 17.16	C	C
	ATOM	4403		THR	159		-44.930		1.00 16.11	С	C
_	MOTA	4404	0	THR	159		-43.710		1.00 17.45	С	0
5	MOTA	4405		HIS	160		-45.742		1.00 15.29	C	N
	MOTA	4406		HIS	160		-45.257		1.00 14.74	С	С
	MOTA	4407	CB	HIS	160	7.397	-46.430	166.912	1.00 14.28	С	C
	MOTA	4408	CG	HIS	160	8.445	-46.055	165.905	1.00 13.23	С	С
	MOTA	4409	CD2	HIS	160	8.627	-46.444	164.620	1.00 13.10	C	С
10	MOTA	4410	ND1	HIS	160	9.462	-45.168	166.182	1.00 13.39	С	N
	ATOM	4411	CE1	HIS	160	10.228	-45.027	165.112	1.00 13.13	С	C
	ATOM	4412	NE2	HIS	160	9.744	-45.792	164.151	1.00 12.34	С	N
	MOTA	4413	С	HIS	160		-44.541		1.00 14.39	C	C
	ATOM	4414	0	HIS	160		-43.463		1.00 14.03	Č	ō
15	ATOM	4415	N	PHE	161		-45.144		1.00 14.40	Č	N
. •	ATOM	4416	CA	PHE	161		-44.526		1.00 15.29	c	C
,		4417	CB	PHE	161		-45.472	_	1.00 15.25	C	C
•	ATOM	4418	CG	PHE	161		-46.510		1.00 13.33		C
		4419	CD1							C	
20	ATOM	_	CD1		161		-46.417		1.00 15.87	C	C
20	MOTA	4420			161		-47.558		1.00 17.11	C	С
	ATOM	4421		PHE	161		-47.341		1.00 17.22	С	С
	MOTA	4422		PHE	161		-48.496		1.00 18.15	C	C
	MOTA	4423	CZ	PHE	161		-48.388		1.00 17.06	С	C
	ATOM	4424	C	PHE	161			164.636	1.00 14.79	C	С
25	MOTA	4425	0	PHE	161			163.928	1.00 13.30	С	0
	MOTA	4426	N	ALA	162	2.954	-43.156	165.732	1.00 15.09	С	N
	MOTA	4427	CA	ALA	162			166.153	1.00 15.95	С	C
	MOTA	4428	CB	ALA	162	1.532	-42.135	167.440	1.00 15.00	C	С
	ATOM	4429	C	ALA	162	3.376	-40.835	166.358	1.00 15.56	С	C
30	ATOM	4430	0	ALA	162	3.162	-39.687	165.986	1.00 15.40	С	0
	ATOM	4431	N	ASP	163	4.521	-41.201	166.938	1.00 16.37	С	N
	ATOM	4432	CA	ASP	163			167.165	1.00 17.11	C	C
	ATOM	4433	CB	ASP	163			168.039	1.00 18.76	С	C
	ATOM	4434	CG	ASP	163			169.518	1.00 19.80	Č	Č
35	ATOM	4435		ASP	163			170.303	1.00 20.29	Ċ	ŏ
•	ATOM	4436		ASP	163			169.896	1.00 20.75	Č	ŏ
	ATOM	4437	C	ASP	163			165.879	1.00 16.41	C	č
	ATOM	4438	Ö	ASP	163			165.703	1.00 16.53	C	0
	ATOM	4439	N	ILE	164			164.976	1.00 15.33	c	N
40	ATOM	4440	CA	ILE	164			163.747			C
40	ATOM	4441	CB	ILE	164			163.747	1.00 14.57	C	
			CG2						1.00 14.61		C
	ATOM	4442			164			163.974		C	
	ATOM	4443		ILE	164			162.497	1.00 13.93	C	C
45	ATOM	4444		ILE	164			161.668	1.00 15.42	С	С
45	ATOM	4445	С	ILE	164			162.801	1.00 14.37	C	C
	ATOM	4446	0	ILE	164			162.025	1.00 13.76	С	0
	MOTA	4447	N	ASN	165			162.854	1.00 13.47	С	N
	ATOM	4448	CA	ASN	165			162.018	1.00 14.22	C	C
	MOTA	4449	CB	ASN	165			162.189	1.00 13.22	С	C
50	ATOM	4450	CG	ASN	165	2.520	-41.304	161.424	1.00 13.80	С	C
	MOTA	4451		ASN	165			160.545	1.00 13.34	С	0
	ATOM	4452	ND2	ASN	165	1.496	-42.089	161.748	1.00 11.89	С	N
	MOTA	4453	С	ASN	165	3.847	-37.809	162.453	1.00 13.40	С	С
	MOTA	4454	0	ASN	165			161.620	1.00 12.46	C	ō
55	MOTA	4455	N	THR	166			163.764	1.00 14.10		N
	MOTA	4456	CA	THR	166			164.302	1.00 15.32		C
	MOTA	4457	СВ	THR	166			165.807	1.00 16.12		Ċ
	ATOM	4458		THR	166			165.995	1.00 15.30		o
		0				203	555	200.975	1.00 13.30	_	9

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	ATOM	4459	CG2	THR	166	3.055	-34.869	166.348	1.00 14.52	С	C
	MOTA	4460	C	THR	166	4.968	-35.403	164.064	1.00 15.86	C	C
	ATOM	4461	0	THR	166	4.903	-34.223	163.724	1.00 16.48	C	0
	MOTA	4462	N	PHE	167	6.124	-36.035	164.234	1.00 17.03	C	N
5	ATOM	4463	CA	PHE	167	7.400	-35.373	163.990	1.00 16.78	Č	C
	MOTA	4464	СВ	PHE	167		-36.363		1.00 17.79	Ċ	Ċ
	ATOM	4465	CG	PHE	167		-35.942		1.00 18.70	Ċ	Ċ
	ATOM	4466	CD1		167		-34.915		1.00 19.29	c	Č
	ATOM	4467	CD2		167		-36.594		1.00 19.25	c	C
10	ATOM	4468		PHE	167		-34.545		1.00 19.64	C	Ċ
. •	ATOM	4469		PHE	167		-36.234		1.00 19.87	C	C
	ATOM	4470	CZ	PHE	167		-35.208		1.00 19.58	c	C
	MOTA	4471	C	PHE	167		-34.887		1.00 15.38	C	C
	ATOM	4472	Ö	PHE	167			162.352	1.00 16.84	C	0
15	ATOM	4473	N	MET	168		-35.755				
ı,J	ATOM	4474	CA	MET	168				1.00 15.69	С	N
	ATOM	4475	CB	MET	168			160.187	1.00 14.91	C	C
								159.312	1.00 14.73	C	C
	MOTA	4476	CG	MET	168			159.161	1.00 12.77	C	C
20	ATOM	4477	SD	MET	168			157.918	1.00 13.46	С	S
20	MOTA	4478	CE	MET	168			158.891	1.00 9.89	С	С
	MOTA	4479	C	MET	168			159.890	1.00 15.70	C	С
	MOTA	4480	0	MET	168			159.097	1.00 13.88	C	0
	MOTA	4481	N	VAL	169			160.510	1.00 16.72	C	N
05	MOTA	4482	CA	VAL	169			160.303	1.00 17.84	С	С
25	MOTA	4483	CB	VAL	169			161.132	1.00 18.74	C	С
	MOTA	4484		VAL	169			161.111	1.00 16.96	С	С
	MOTA	4485		VAL	169			160.544	1.00 17.94	C	С
	MOTA	4486	С	VAL	169			160.694	1.00 18.62	С	С
	MOTA	4487	0	VAL	169			159.954	1.00 18.14	С	0
30	MOTA	4488	N	LEU	170			161.850	1.00 19.05	С	N
	MOTA	4489	CA	LEU	170			162.284	1.00 18.80	С	С
	MOTA	4490	CB	LEU	170			163.700	1.00 19.18	C	С
	MOTA	4491	CG	LEU	170			164.808	1.00 20.13	C	C
	MOTA	4492		LEU	170			166.112	1.00 20.19	C	C
35	ATOM	4493		LEU	170	4.560	-29.665	164.983	1.00 21.80	C	С
	MOTA	4494	С	LEU	170			161.297	1.00 19.15	С	C
	MOTA	4495	0	LEU	170	7.167	-28.989	161.055	1.00 19.95	C	0
	MOTA	4496	N	GLN	171	7.670	-31.148	160.718	1.00 18.11	C	N
	ATOM	4497	CA	GLN	171	8.713	-30.819	159.743	1.00 17.90	С	С
40	ATOM	4498	CB	GLN	171	9.557	-32.050	159.382	1.00 17.69	С	С
	MOTA	4499	CG	GLN	171	10.492	-32.522	160.501	1.00 19.25	С	С
	ATOM	4500	CD	GLN	171	11.383	-31.408	161.040	1.00 19.45	С	
	ATOM	4501	OE1	GLN	171	12.164	-30.802	160.305	1.00 21.44	С	0
	ATOM	4502		GLN	171			162.328	1.00 19.42	Ċ	N
45	ATOM	4503	С	GLN	171			158.458	1.00 17.82	Ċ	C
	ATOM	4504	0	GLN	171			157.841	1.00 17.25	Ċ	ō
	ATOM	4505	N	VAL	172			158.039	1.00 17.26	Č	N
	ATOM	4506	CA	VAL	172			156.834	1.00 18.43	č	C
	ATOM	4507	СВ	VAL	172			156.376	1.00 18.66	Č	Č
50	ATOM	4508		VAL	172			155.185	1.00 17.80	Č	c
	ATOM	4509		VAL	172			155.967	1.00 18.65	Č	C
	ATOM	4510	C	VAL	172			157.107	1.00 18.42	C	C
	MOTA	4511	Ö	VAL	172			156.264	1.00 18.42	C	
	ATOM	4512	N	ILE	173			158.286			0
55	ATOM	4513	CA	ILE	173			158.286	1.00 18.42	C	N
<b>J</b> J	MOTA	4513	CB	ILE	173			160.090	1.00 18.94	C	C
		4514		ILE	173				1.00 19.45	C	C
	ATOM				173			160.550	1.00 18.53	C	C
	MOTA	4516	CGI	ILE	1/3	4.9/6	-21.186	160.147	1.00 19.67	С	С

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	MOTA	4517	CD1		173		-28.048		1.00 18.69	С	C
	MOTA	4518	С	ILE	173	6.077	-26.128	158.509	1.00 18.47	C	С
	MOTA	4519	0	ILE	173		-25.093		1.00 17.88	С	0
_	MOTA	4520	N	LYS	174	7.209	-26.498	159.103	1.00 17.67	С	N
5	MOTA	4521	CA	LYS	174		-25.671		1.00 18.85	С	С
	MOTA	4522	CB	LYS	174	9.559	-26.325	159.797	1.00 19.84	C	С
	MOTA	4523	CG	LYS	174	9.282	-26.514	161.293	1.00 20.37	С	С
	MOTA	4524	CD	LYS	174		-27.203		1.00 22.31	С	С
	MOTA	4525	CE	LYS	174		-26.287		1.00 24.43	C	С
10	MOTA	4526	NZ	LYS	174		-26.910		1.00 27.02	С	N
	MOTA	4527	С	LYS	174	8.826	-25.447	157.568	1.00 19.16	С	C
	MOTA	4528	0	LYS	174			157.194	1.00 18.69	C	0
	MOTA	4529	N	PHE	175		-26.499		1.00 18.61	С	N
	MOTA	4530	CA	PHE	175		-26.432		1.00 18.87	С	С
15	MOTA	4531	СВ	PHE	175			154.697	1.00 18.31	С	С
	MOTA	4532	CG	PHE	175			153.190	1.00 18.11	С	C
	MOTA	4533		PHE	175			152.485	1.00 16.75	С	С
	MOTA	4534		PHE	175	7.702	-28.099	152.478	1.00 17.31	C	С
	MOTA	4535		PHE	175			151.090	1.00 17.12	C	C
20	MOTA	4536	CE2	PHE	175			151.079	1.00 16.84	С	С
	MOTA	4537	CZ	PHE	175	8.854	-27.742	150.386	1.00 16.17	С	С
	MOTA	4538	С	PHE	175			154.619	1.00 19.59	C	С
	MOTA	4539	0	PHE	175			153.875	1.00 19.63	C	0
	MOTA	4540	N	THR	176			154.834	1.00 19.92	C	N
25	MOTA	4541	CA	THR	176			154.181	1.00 21.48	C	C
	ATOM	4542	CB	THR	176			154.427	1.00 21.69	C	С
	MOTA	4543	OG1	THR	176	4.209	-24.860	155.833	1.00 22.61	C	0
	ATOM	4544	CG2		176			153.886	1.00 20.21	C	С
	MOTA	4545	С	THR	176			154.668	1.00 22.48	С	C
30	ATOM	4546	0	THR	176			153.896	1.00 22.19	C	0
	ATOM	4547	N	LYS	177			155.946	1.00 22.93	C	N
	MOTA	4548	CA	LYS	177			156.491	1.00 24.72	С	C
	MOTA	4549	СВ	LYS	177			158.018	1.00 25.60	С	С
	MOTA	4550	CG	LYS	177			158.658	1.00 28.46	С	С
35	MOTA	4551	CD	LYS	177			160.169	1.00 28.76	С	С
	MOTA	4552	CE	LYS	177			160.754	1.00 28.66	С	С
	MOTA	4553	NZ	LYS	177			160.208	1.00 29.61	С	N
	MOTA	4554	C	LYS	177			155.900	1.00 24.56	С	С
40	MOTA	4555	0	LYS	177			155.971	1.00 23.75	C	0
40	ATOM	4556	N	ASP	178			155.313	1.00 24.31	С	N
	MOTA	4557	CA	ASP	178			154.684	1.00 25.00	С	С
	MOTA	4558	CB	ASP	178			154.849	1.00 24.90	С	
	ATOM	4559	CG	ASP	178			156.238	1.00 25.48	С	С
45	ATOM	4560		ASP	178			157.090	1.00 26.26	C	0
45	MOTA	4561		ASP	178			156.478	1.00 24.33	С	0
	MOTA	4562	C	ASP	178			153.199	1.00 25.18	C	C
	MOTA	4563	0	ASP	178			152.448	1.00 26.75	С	0
	MOTA	4564	N	LEU	179			152.772	1.00 24.75	C	N
50	ATOM	4565	CA	LEU	179			151.379	1.00 24.20	С	C
50	ATOM	4566	CB	LEU	179			150.822	1.00 23.10	С	С
	MOTA	4567	CG	LEU	179			150.875	1.00 22.16	С	С
	MOTA	4568		LEU	179			150.183	1.00 21.20	C	С
	MOTA	4569		LEU	179			150.213	1.00 20.84	С	C
	MOTA	4570	C	LEU	179			151.264	1.00 24.41	C	C
55	ATOM	4571	0	LEU	179			151.623	1.00 24.15	C	0
	ATOM	4572	N	PRO	180			150.776		C	N
	ATOM	4573	CD	PRO	180			150.417		С	С
	MOTA	4574	CA	PRO	180	7.456	-16.886	150.627	1.00 25.07	С	С

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	ATOM	4575	СВ	PRO	180	8.446 -16.102 149.771 1.00 25.45	C
	ATOM	4576	CG	PRO	180	9.757 -16.576 150.309 1.00 25.01	
	ATOM	4577	C	PRO	180	6.062 -16.973 150.003 1.00 25.06	
	MOTA	4578	0	PRO	180	5.107 -16.419 150.543 1.00 24.75	
5		4579	-				
9	MOTA		N	VAL	181	5.937 -17.676 148.880 1.00 25.37 (	
	MOTA	4580	CA	VAL	181	4.637 -17.808 148.218 1.00 26.24 (	
	MOTA	4581	CB	VAL	181	4.743 -18.624 146.898 1.00 27.68 (	
	MOTA	4582	CG1		181	5.515 -17.822 145.857 1.00 28.55	
40	MOTA	4583	CG2		181		C
10	MOTA	4584	С	VAL	181		C C
	MOTA	4585	0	VAL	181		0
	MOTA	4586	N	PHE	182		CN
	MOTA	4587	CA	PHE	182	3.062 -19.988 150.933 1.00 25.66	C C
	MOTA	4588	СВ	PHE	182		C C
15	ATOM	4589	CG	PHE	182	2.859 -21.923 152.598 1.00 24.52	C C
	MOTA	4590	CD1	PHE	182		СС
	MOTA	4591	CD2	PHE	182		СС
	MOTA	4592	CE1	PHE	182		c c
	MOTA	4593	CE2	PHE	182		СС
20	MOTA	4594	CZ	PHE	182		c · c
	MOTA	4595	C	PHE	182		c c
	ATOM	4596	Ō	PHE	182		c o
	ATOM	4597	N	ARG	183		CN
	ATOM	4598	CA	ARG	183		CC
25	ATOM	4599	CB	ARG	183		C C
	ATOM	4600	CG	ARG	183		c c
	MOTA	4601	CD	ARG	183		CC
	MOTA	4602	NE	ARG	183	•	C N
	ATOM	4603	CZ	ARG	183		CC
30	MOTA	4604		ARG	183		CN
00	ATOM	4605	NH2		183		C N
	MOTA	4606	C	ARG	183		CC
	ATOM	4607	Ö	ARG	183		
	ATOM	4608	N	SER	184		
35	ATOM	4608	CA	SER			C N
33					184		C C
	MOTA	4610	CB	SER	184		C C
	MOTA	4611	OG	SER	184		C O
	MOTA	4612	C	SER	184		C C
40	MOTA	4613	0	SER	184		C O
40	MOTA	4614	N	LEU	185		C N
	ATOM	4615	CA	LEU	185		C C
	ATOM	4616	CB	LEU	185		C C
	MOTA	4617	CG	LEU	185		C C
4.5	MOTA	4618		LEU	185		C C
45	MOTA	4619		LEU	185		C C
	MOTA	4620	С	LEU	185		C C
	MOTA	4621	0	LEU	185		C O
	MOTA	4622	N	PRO	186	-3.455 -16.085 152.703 1.00 33.52	C N
	MOTA	4623	CD	PRO	186		C C
50	MOTA	4624	CA	PRO	186		C C
	MOTA	4625	CB	PRO	186	-5.564 -15.552 153.588 1.00 34.21	C C
	MOTA	4626	CG	PRO	186	-5.714 -16.243 152.264 1.00 34.41	C C
	MOTA	4627	С	PRO	186	-3.952 -17.050 154.880 1.00 35.52	C C
	MOTA	4628	0	PRO	186	-3.863 -18.176 154.389 1.00 35.54	C O
55	MOTA	4629	N	ILE	187	-3.907 -16.831 156.189 1.00 36.77	C N
	ATOM	4630	CA	ILE	187	-3.728 -17.921 157.143 1.00 37.62	c c
	ATOM	4631	СВ	ILE	187	-3.844 -17.417 158.594 1.00 38.59	c c
	MOTA	4632	CG2		187	-3.717 -18.588 159.562 1.00 39.09	CC

	MOTA	4633	CG1 IL	E 187		-16.381		1.00 39.17	С	С
	ATOM	4634	CD1 IL	E 187	-1.337	-16.901	158.653	1.00 39.06	С	С
	MOTA	4635	C IL	E 187	-4.682	-19.092	156.964	1.00 37.72	C	С
_	ATOM	4636	O IL	E 187	-4.264	-20.246	157.055	1.00 37.65	С	0
5	MOTA	4637	N GL	U 188	-5.959	-18.808	156.717	1.00 37.53	C	N
	MOTA	4638	CA GL	U 188	-6.939	-19.878	156.535	1.00 37.86	С	С
	ATOM	4639	CB GL	U 188	-8.341	-19.302	156.286	1.00 39.84	С	С
	MOTA	4640	CG GL	U 188	-8.424	-18.286	155.155	1.00 43.27	С	С
	ATOM	4641	CD GL	U 188	-8.379	-16.849	155.645	1.00 45.23	С	C
10	ATOM	4642	OE1 GL	U 188	-7.596	-16.550	156.578	1.00 45.96	С	0
	ATOM	4643	OE2 GI	U 188	-9.125	-16.015	155.084	1.00 46.68	С	0
	ATOM	4644	C GI		-6.558	-20.832	155.397	1.00 36.71	С	C
	MOTA	4645	O GL	U 188	-6.786	-22.041	155.487	1.00 35.74	С	0
	MOTA	4646	N AS	P 189	-5.977	-20.292	154.330	1.00 35.44	C	N
15	MOTA	4647	CA AS	P 189	-5.563	-21.114	153.199	1.00 34.24	С	С
	MOTA	4648	CB AS	P 189	-5.378	-20.255	151.946	1.00 35.42	С	С
	MOTA	4649	CG AS	P 189	-6.701	-19.753	151.380	1.00 36.94	С	С
	MOTA	4650	OD1 AS	P 189	-6.722	-19.324	150.210	1.00 38.00	С	0
	MOTA	4651	OD2 AS	SP 189	-7.717	-19.781	152.105	1.00 38.31	С	0
20	MOTA	4652	C AS	SP 189	-4.279	-21.891	153.503	1.00 32.76	С	С
	MOTA	4653	O AS	SP 189	-4.089	-22.998	153.001	1.00 31.31	С	0
	MOTA	4654	N GI	N 190	-3.397	-21.311	154.313	1.00 31.50	С	N
	MOTA	4655	CA GI	N 190	-2.165	-21.995	154.694	1.00 31.34	С	С
	MOTA	4656	CB GI	N 190	-1.279	-21.090	155.565	1.00 31.81	С	С
25	MOTA	4657	CG GI	N 190	-0.838	-19.803	154.884	1.00 32.88	С	С
	ATOM	4658	CD GI	N 190	0.168	-19.006	155.699	1.00 33.88	C	C
	MOTA	4659	OE1 GI	LN 190	0.376	-17.816	155.452	1.00 35.31	С	0
	MOTA	4660	NE2 GI	LN 190	0.805	-19.658	156.663	1.00 32.82	C	N
	ATOM	4661	C GI	LN 190	-2.569	-23.242	155.484	1.00 30.49	С	C
30	MOTA	4662	O G1	LN 190	-2.039	-24.326	155.258	1.00 29.34	С	0
	ATOM	4663	N II	LE 191	-3.520	-23.070	156.402	1.00 30.50	С	N
	MOTA	4664	CA I	LE 191	-4.036	-24.163	157.227	1.00 30.60	С	С
	ATOM	4665	CB I	LE 191	-5.216	-23.701	158.133	1.00 31.67	С	С
	ATOM	4666	CG2 I	LE 191	-5.683	-24.857	159.000	1.00 30.87	С	C
35	ATOM	4667	CG1 I	LE 191	-4.805	-22.512	159.006	1.00 32.04	С	C
	MOTA	4668	CD1 I	LE 191	-3.723	-22.813	159.995	1.00 33.57	С	С
	ATOM	4669	C I	LE 191	-4.574	-25.275	156.334	1.00 30.25	C	С
	ATOM	4670	O I	LE 191	-4.203	-26.445	156.476	1.00 29.94	С	0
	ATOM	4671	N S	ER 192	-5.458	-24.899	155.415	1.00 28.90	С	N
40	MOTA	4672	CA S	ER 192	-6.075	-25.853	154.503	1.00 28.72	С	С
	ATOM	4673	CB S	ER 192	-7.119	-25.147	153.644	1.00 28.55	C	C
	ATOM	4674	OG S	ER 192	-8.087	-24.534	154.472	1.00 31.09	С	0
	MOTA	4675	C S	ER .192	-5.087	-26.587	153.605	1.00 27.78	С	C
	MOTA	4676	0 S	ER 192	-5.200	-27.797	153.420	1.00 27.73	С	0
45	ATOM	4677	N L	EU 193	-4.129	-25.867	153.033	1.00 26.80	C	N
	ATOM	4678	CA L	EU 193	-3.158	-26.519	152.168	1.00 26.77	С	С
	MOTA	4679	CB L	EU 193	-2.310	-25.489	151.415	1.00 26.32	С	C
	ATOM	4680	CG L	EU 193	-3.046	-24.625	150.384	1.00 26.16	С	С
	MOTA	4681	CD1 L	EU 193	-2.009	-23.922	149.513	1.00 26.75	С	C
50	ATOM	4682	CD2 L	EU 193	-3.969	-25.479	149.519	1.00 24.72	С	С
	ATOM	4683	C L	EU 193			152.952	1.00 26.50	С	C
	ATOM	4684		EU 193			152.468	1.00 25.42	C	Ō
	ATOM	4685		EU 194			154.159	1.00 26.75	C	N
	ATOM	4686		EU 194			154.991	1.00 27.59	C	C
55	ATOM	4687		EU 194			156.301		С	C
	ATOM	4688		EU 194			157.201		C	Č
	ATOM	4689	CD1 L				156.797		C	Ċ
	ATOM	4690	CD2 L				158.662		C	Ċ
					,					

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	MOTA	4691	С	LEU	194	-1.717	-29.246	155.323	1.00 27.29	С	С
	MOTA	4692	0	LEU	194	-1.151	-30.328	155.139	1.00 26.70	C	0
	MOTA	4693	N	LYS	195	-2.948	-29.148	155.818	1.00 26.53	С	N
	ATOM	4694	CA	LYS	195			156.172	1.00 27.06	C	С
5	ATOM	4695		LYS	195		-29.918		1.00 28.79	Ċ	C
	ATOM	4696		LYS	195			158.138	1.00 31.86	Č	Ċ
	ATOM	4697	CD	LYS	195			158.740	1.00 34.00	č	Ċ
	ATOM	4698	CE	LYS	195		-28.259		1.00 35.25	Č	Ċ
	ATOM	4699	NZ	LYS	195			160.781	1.00 37.11	č	Ŋ
10	ATOM	4700	C	LYS	195			154.956	1.00 26.13	č	C
	ATOM	4701	0	LYS	195			155.048	1.00 26.21	č	ŏ
	ATOM	4702	N	GLY	196			153.812	1.00 24.83	c	N
	ATOM	4703	CA	GLY	196			152.606	1.00 23.63	c	C
	ATOM	4704	C	GLY	196			151.969	1.00 22.89	č	c
15	ATOM	4705	Ö	GLY	196			151.306	1.00 21.84	C	Ö
	ATOM	4706	N	ALA	197			152.179	1.00 21.49	C	N
	ATOM	4707	CA	ALA	197			151.545	1.00 21.43	C	C
	ATOM	4708	СВ	ALA	197			150.584	1.00 21.24	C	C
	ATOM	4709	C	ALA	197			152.421	1.00 19.04	C	C
20	MOTA	4710	0	ALA	197			152.421			
20	ATOM	4711	N	ALA	198			151.905		C	0
	ATOM	4712	CA	ALA	198				1.00 18.80	C	N
		4712						154.616	1.00 17.78	C	C
	MOTA		CB	ALA	198			156.052	1.00 17.25	C	C
25	ATOM	4714	C	ALA	198			154.448	1.00 17.03	C	C
25	MOTA	4715	0	ALA	198			154.304	1.00 16.38	C	0
	ATOM	4716	N	VAL	199			154.482	1.00 16.43	C	N
	MOTA	4717	CA	VAL	199			154.332	1.00 16.70	C	C
	ATOM	4718	CB	VAL	199			154.613	1.00 16.81	С	С
20	ATOM	4719		VAL	199			154.262	1.00 17.35	С	C
30	MOTA	4720		VAL	199			156.093	1.00 16.94	С	С
	ATOM	4721	C	VAL	199			152.934	1.00 16.67	С	С
	MOTA	4722	0	VAL	199			152.778	1.00 15.71	С	0
	ATOM	4723	N	GLU	200			151.920	1.00 16.31	C	N
0.0	MOTA	4724	CA	GLU	200			150.546	1.00 17.25	С	C
35	MOTA	4725	СВ	GLU	200			149.589	1.00 18.45	С	С
	MOTA	4726	CG	GLU	200			149.242	1.00 19.72	С	С
	MOTA	4727	CD	GLU	200			148.399	1.00 20.49	С	C
	MOTA	4728		GLU	200			147.526	1.00 20.99	С	0
	ATOM	4729		GLU	200			148.601	1.00 21.11	С	0
40	MOTA	4730	С	GLU	200			150.373	1.00 17.44	С	C
	ATOM	4731	0	GLU	200			149.792	1.00 16.76	C	0
	MOTA	4732	N	ILE	201			150.875	1.00 17.66	С	N
	ATOM	4733	CA	ILE	201			150.794	1.00 17.75	С	С
	MOTA	4734	СВ	ILE	201			151.458	1.00 18.15	С	С
45	ATOM	4735		ILE	201	6.057	-32.649	151.706	1.00 17.40	С	С
	MOTA	4736	CG1	ILE	201	3.885	-32.018	150.561	1.00 17.68	С	C
	MOTA	4737	CD1	ILE	201	3.686	-30.658	151.190	1.00 15.43	С	C
	MOTA	4738	С	ILE	201			151.491	1.00 18.18	С	С
	MOTA	4739	0	ILE	201	6.542	-35.723	150.976	1.00 18.01	С	0
50	ATOM	4740	N	CYS	202	5.077	-35.890	152.669	1.00 17.95	С	N
	MOTA	4741	CA	CYS	202	5.917	-36.812	153.417	1.00 18.27	С	С
	ATOM	4742	CB	CYS	202	5.259	-37.161	. 154.752	1.00 17.78	С	
	ATOM	4743	SG	CYS	202			155.941	1.00 19.31	C	S
	ATOM	4744	С	CYS	202			152.632	1.00 17.48	С	c s c
55	ATOM	4745	0	CYS	202			152.684	1.00 17.82	C	ō
	MOTA	4746	N	HIS	203			151.908	1.00 16.38	Č	N
	ATOM	4747	CA	HIS	203			151.121	1.00 16.70	Č	Ċ
	MOTA	4748	CB	HIS	203			150.611	1.00 15.92	Č	Č
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	MOTA	4749		HIS	203		-41.272		1.00 15.86	C	С
	MOTA	4750	CD2		203		-41.080		1.00 16.22	C	С
	MOTA	4751	ND1		203		-42.439		1.00 16.53	С	N
_	MOTA	4752	CE1		203		-42.930		1.00 16.84	С	С
5	MOTA	4753	NE2		203		-42.123		1.00 16.72	C	N
	MOTA	4754		HIS	203		-39.519		1.00 16.55	С	С
	MOTA	4755	0	HIS	203		-40.352		1.00 17.66	С	0
	ATOM	4756		ILE	204		-38.331		1.00 16.27	C	N
40	MOTA	4757		ILE	204		-37.944		1.00 16.29	C	С
10	MOTA	4758		ILE	204		-36.571		1.00 16.08	С	С
	MOTA	4759		ILE	204		-36.055	_	1.00 16.01	C	С
	MOTA	4760		ILE	204		-36.708		1.00 14.17	С	С
	MOTA	4761		ILE	204		-35.396		1.00 13.55	С	C
45	MOTA	4762		ILE	204		-37.875		1.00 17.25	С	С
15	MOTA	4763		ILE	204		-38.383		1.00 17.11	C	0
	MOTA	4764		VAL	205		-37.248		1.00 17.08	С	N
	MOTA	4765		VAL	205		-37.121		1.00 18.01	С	С
	ATOM	4766		VAL	205		-36.200		1.00 17.13	С	С
20	ATOM	4767	CG1		205		-36.276		1.00 16.96	С	C
20	MOTA	4768	CG2		205		-34.765		1.00 16.00	C	C
	ATOM	4769		VAL	205		-38.486		1.00 18.51	C	C
	ATOM	4770		VAL	205		-38.765		1.00 19.15	C	0
	ATOM	4771	N	LEU	206		-39.331		1.00 19.05	C	N
25	MOTA	4772	CA	LEU	206		-40.671		1.00 20.76	C	C
25	MOTA	4773 4774	CB CG	LEU	206		-41.285 -41.352		1.00 21.98	C	C
	MOTA	4775		LEU	206				1.00 24.68	C	C
	MOTA	4776	CD1 CD2		206			154.882	1.00 24.95	C	C
	MOTA MOTA	4777	CDZ	LEU	206 206			155.032 150.841	1.00 24.49 1.00 20.80	C	C C
30	ATOM	4778	0	LEU	206			151.055	1.00 20.80	C	
30	ATOM	4779	N	ASN	207			149.623	1.00 20.84	C	0
	ATOM	4780	CA	ASN	207			149.623	1.00 19.96	C	N C
	ATOM	4781	CB	ASN	207			147.199	1.00 20.37	C	C
	MOTA	4782	CG	ASN	207			146.017	1.00 18.23	C	C
35	ATOM	4783	OD1		207		-42.259		1.00 18.37	C	0
00	ATOM	4784	ND2		207			146.123	1.00 16.39	C	N
	ATOM	4785	C	ASN	207			148.280	1.00 10.71		C
	ATOM	4786	Ö	ASN	207			147.885	1.00 20.03	C	Ö
	ATOM	4787	N	THR	208			148.556	1.00 20.81		N
40	ATOM	4788	CA	THR	208			148.403	1.00 22.30		C
	ATOM	4789	СВ	THR	208			148.528	1.00 22.75		c
	MOTA	4790		THR	208			149.624	1.00 22.99	Č	ŏ
	MOTA	4791		THR	208			147.239	1.00 25.40		Č
	MOTA	4792	С	THR	208			149.372	1.00 21.11		C
45	MOTA	4793	0	THR	208			149.190	1.00 21.58		Ō
	ATOM	4794	N	THR	209	14.061	-43.095	150.400	1.00 20.44		N
	MOTA	4795	CA	THR	209			151.308	1.00 20.16		C
	MOTA	4796	СВ	THR	209			152.773	1.00 21.00		C
	ATOM	4797	OG1	THR	209			152.920	1.00 20.06		0
50	ATOM	4798	CG2	THR	209	14.310	-42.427	153.178	1.00 19.21		С
	MOTA	4799	С	THR	209	14.086	-45.511	150.869	1.00 20.37		C
	ATOM	4800	0	THR	209	14.535	-46.528	151.402	1.00 19.73		0
	ATOM	4801	N	PHE	210			149.886	1.00 19.78		N
	MOTA	4802	CA	PHE	210	12.623	-46.803	149.404	1.00 21.06		C
55	MOTA	4803	CB	PHE	210			148.552	1.00 20.74	C	С
	MOTA	4804	CG	PHE	210			148.271	1.00 20.23		C
	MOTA	4805		PHE	210			149.303	1.00 20.70		C
	MOTA	4806	CD2	PHE	210	10.392	-48.204	146.969	1.00 19.93	C	C

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	MOTA	4807	CE1	PHE	210	9.062	-49.491	149.045	1.00 21.07	С	С
	MOTA	4808	CE2	PHE	210	9.592	-49.310	146.701	1.00 19.80	C	С
	MOTA	4809	CZ	PHE	210	8.924	-49.955	147.740	1.00 20.40	C	С
_	ATOM	4810	С	PHE	210	13.605	-47.647	148.602	1.00 21.28	C	С
5	MOTA	4811	0	PHE	210	14.202	-47.186	147.640	1.00 20.88	C	0
	MOTA	4812	N	CYS	211	13.770	-48.892	149.015	1.00 22.28	С	N
	MOTA	4813	CA	CYS	211	14.670	-49.802	148.330	1.00 24.70	C	C
	MOTA	4814	CB	CYS	211	15.421	-50.656	149.352	1.00 25.33	С	С
	MOTA	4815	SG	CYS	211	16.567	-51.838	148.615	1.00 28.07	С	S
10	ATOM	4816	С	CYS	211	13.834	-50.694	147.417	1.00 25.31	С	С
	ATOM	4817	0	CYS	211	13.077	-51.533	147.889	1.00 24.53	С	0
	ATOM	4818	N	LEU	212	13.964	-50.502	146.111	1.00 26.95	С	N
	ATOM	4819	CA	LEU	212	13.206	-51.290	145.142	1.00 29.54	C	С
	ATOM	4820	CB	LEU	212	13.581	-50.859	143.724	1.00 29.30	С	С
15	ATOM	4821	CG	LEU	212	13.098	-49.465	143.324	1.00 30.74	C	С
	ATOM	4822	CD1	LEU	212	13.746	-49.044	142.004	1.00 30.10	С	С
	ATOM	4823	CD2	LEU	212		-49.475		1.00 28.92	С	С
	MOTA	4824	С	LEU	212		-52.795		1.00 30.81	С	С
	ATOM	4825	0	LEU	212		-53.587		1.00 30.78	C	0
20	ATOM	4826	N	GLN	213			145.469	1.00 31.69	Ċ	N
	ATOM	4827	CA	GLN	213			145.616	1.00 33.12	Č	C
	ATOM	4828	CB	GLN	213			145.725	1.00 35.87	Č	C
	ATOM	4829	CG	GLN	213			145.991	1.00 39.28	Č	Č
	ATOM	4830	CD	GLN	213			146.143	1.00 41.77	Č	Ċ
25	ATOM	4831		GLN	213			146.398	1.00 43.80	Č	ō
	MOTA	4832	NE2		213			145.985	1.00 43.06	Č	N
	ATOM	4833	С	GLN	213			146.792	1.00 32.23	Č	C
	ATOM	4834	Ō	GLN	213			146.667	1.00 32.37	Č	ō
	ATOM	4835	N	THR	214			147.930	1.00 30.26	C	N
30	ATOM	4836	CA	THR	214			149.103	1.00 28.07	C	C
•	ATOM	4837	CB	THR	214			150.331	1.00 28.29	C	C
	ATOM	4838	OG1		214			150.645	1.00 27.85	C	0
	ATOM	4839	CG2		214			150.047	1.00 28.08	C	C
	ATOM	4840	C	THR	214			149.509	1.00 27.47	c	C
35	ATOM	4841	Õ	THR	214			150.420	1.00 27.47	C	0
00	ATOM	4842	N	GLN	215			148.844	1.00 26.24	C	
	MOTA	4843	CA	GLN	215			149.175	1.00 20.24	c	N C
	MOTA	4844	CB	GLN	215			148.961	1.00 27.14	C	С
	ATOM	4845	CG	GLN	215			147.533	1.00 28.32	C	C
40	ATOM	4846	CD	GLN	215			146.532	1.00 32.89		
40	ATOM	4847		GLN	215			146.532		C	C
	ATOM	4848		GLN	215			145.509	1.00 36.92	C	0
	ATOM	4849	C	GLN	215			150.639	1.00 35.80	C	N
								151.318	1.00 25.79	C	C
45	ATOM	4850	0	GLN	215				1.00 25.44	C	0
45	ATOM	4851	N	ASN	216			151.103	1.00 24.70	C	N
	ATOM	4852	CA	ASN	216			152.467	1.00 24.04	C	C
	MOTA	4853	CB	ASN	216			153.111	1.00 25.43	C	C
	MOTA	4854	CG	ASN	216			153.744	1.00 28.21	C	C
50	ATOM	4855		ASN	216			153.156	1.00 28.27	C	0
30	ATOM	4856		ASN	216			154.958	1.00 30.94	C	N
	ATOM	4857	C	ASN	216			152.406	1.00 21.89	C	C
	ATOM	4858	0	ASN	216			151.423	1.00 21.17	C	0
	ATOM	4859	N	PHE	217			153.460	1.00 20.68	С	Ŋ
C C	ATOM	4860	CA	PHE	217			153.538	1.00 20.49	С	С
55	MOTA	4861	СВ	PHE	217			154.171	1.00 18.98	С	С
	MOTA	4862	CG	PHE	217			153.268	1.00 18.24	С	C
	MOTA	4863		PHE	217			153.209	1.00 18.43	С	С
	MOTA	4864	CD2	PHE	217	9.728	-45.921	152.472	1.00 19.04	С	C

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	MOTA	4865	CE1		217			152.366	1.00 20.17	C	С
	MOTA	4866		PHE	217			151.622	1.00 18.75	С	С
	MOTA	4867	CZ	PHE	217			151.571	1.00 18.41	C	C
_	MOTA	4868	С	PHE	217			154.472	1.00 20.87	С	С
5	MOTA	4869	0	PHE	217			155.602	1.00 19.76	С	0
	MOTA	4870	N	LEU	218			153.996	1.00 21.69	C	N
	MOTA	4871	CA	LEU	218			154.790	1.00 22.93	C	С
	MOTA	4872	CB	LEU	218			153.910	1.00 24.83	С	С
	MOTA	4873	CG	LEU	218			153.445	1.00 26.67	С	С
10	MOTA	4874	CD1		218			152.686	1.00 28.70	С	С
	MOTA	4875	CD2		218			152.547	1.00 27.72	C	C
	MOTA	4876	С	LEU	218			155.429	1.00 23.17	С	С
	MOTA	4877	0	LEU	218			154.760	1.00 22.44	С	0
	MOTA	4878	N	CYS	219			156.732	1.00 23.14	С	N
15	ATOM	4879	CA	CYS	219			157.435	1.00 24.43	C	C
	MOTA	4880	CB	CYS	219			158.052	1.00 23.99	C	С
	MOTA	4881	SG	CYS	219	13.158	-44.351	156.826	1.00 21.86	C	S
	MOTA	4882	С	CYS	219			158.503	1.00 24.72	С	C
	MOTA	4883	0	CYS	219	16.666	-45.328	159.651	1.00 24.29	С	0
20	MOTA	4884	N	GLY	220	18.181	-44.675	158.105	1.00 24.82	С	N
	MOTA	4885	CA	GLY	220			159.011	1.00 24.67	С	С
	MOTA	4886	С	GLY	220			159.323	1.00 23.88	C	С
	ATOM	4887	0	GLY	220	19.472	-47.078	158.409	1.00 23.43	C	0
	MOTA	4888	N	PRO	221			160.599	1.00 23.24	С	N
25	MOTA	4889	CD	PRO	221			161.801	1.00 23.12	C	C
	MOTA	4890	CA	PRO	221			160.913	1.00 22.65	С	С
	MOTA	4891	CB	PRO	221	20.239	-48.091	162.272	1.00 22.04	С	C
	MOTA	4892	CG	PRO	221	19.635	-46.863	162.926	1.00 21.62	С	C
	MOTA	4893	С	PRO	221			160.964	1.00 22.50	С	С
30	MOTA	4894	0	PRO	221			161.265	1.00 22.01	С	0
	MOTA	4895	N	LEU	222			160.671	1.00 21.70	С	N
	ATOM	4896	CA	LEU	222			160.721	1.00 21.85	С	C
	MOTA	4897	CB	LEU	222			161.316	1.00 21.57	С	С
	MOTA	4898	CG	LEU	222			162.700	1.00 20.39	С	C
35	MOTA	4899		LEU	222			163.131	1.00 20.86	C	C
	MOTA	4900		LEU	222			163.712	1.00 20.79	С	C
	ATOM	4901	С	LEU	222			159.363	1.00 21.95	С	C
	MOTA	4902	0	LEU	222			158.316	1.00 22.14	C	0
	MOTA	4903	N	ARG	223			159.402	1.00 21.42	С	N
40	MOTA	4904	CA	ARG	223			158.198	1.00 21.84	C	C
	MOTA	4905	CB	ARG	223			157.921	1.00 23.26	C	С
	MOTA	4906	CG	ARG	223			157.819	1.00 27.14	C	•
	MOTA	4907	CD	ARG	223			156.482	1.00 30.23	С	С
	MOTA	4908	NE	ARG	223			156.393	1.00 32.56	С	N
45	MOTA	4909	$\mathbf{cz}$	ARG	223			156.860	1.00 33.77	С	C
	MOTA	4910		ARG	223			157.455	1.00 31.71	С	N
	ATOM	4911		ARG	223			156.728	1.00 35.18	С	N
	MOTA	4912	С	ARG	223			158.381	1.00 20.01	С	C
	MOTA	4913	0	ARG	223			159.290	1.00 20.43	C	0
50	MOTA	4914	N	TYR	224			157.517	1.00 17.42	С	N
	MOTA	4915	CA	TYR	224			157.599	1.00 15.88	С	С
	MOTA	4916	CB	TYR	224			157.555	1.00 14.70	C	C
	MOTA	4917	CG	TYR	224			158.674	1.00 15.19	С	С
	MOTA	4918		TYR	224			. 158.488	1.00 14.65	С	С
55	MOTA	4919		TYR	224			159.514	1.00 15.25	С	С
	MOTA	4920		TYR	224			159.924	1.00 14.47	С	C
	MOTA	4921		TYR	224			160.965	1.00 14.99	С	С
	MOTA	4922	CZ	TYR	224	10.615	-46.430	160.746	1.00 14.53	С	С

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	MOTA	4923	ОН	TYR	224		-45.570		1.00 13.71	С	0
	MOTA	4924	С	TYR	224		-51.312		1.00 15.26	С	С
	MOTA	4925	0	TYR	224		-51.087		1.00 12.96	С	0
_	MOTA	4926	N	THR	225		-52.277		1.00 14.87	C	N
5	MOTA	4927	CA	THR	225		-53.162		1.00 15.88	С	С
	MOTA	4928	СВ	THR	225		-54.639		1.00 15.32	C	C
	MOTA	4929	OG1		225		-55.067		1.00 15.98	С	0
	MOTA	4930	CG2	THR	225		-54.799		1.00 15.24	С	С
40	MOTA	4931	С	THR	225		-53.068		1.00 14.70	С	С
10	MOTA	4932	0	THR	225		-52.431		1.00 13.09	C	0
	MOTA	4933	N	ILE	226		-53.727		1.00 14.59	C	N
	MOTA	4934	CA	ILE	226		-53.695		1.00 14.58	С	С
	ATOM	4935	СВ	ILE	226		-54.358		1.00 15.11	C	С
4.5	ATOM	4936		ILE	226		-55.877		1.00 12.89	C	С
15	MOTA	4937	CG1		226		-53.929		1.00 14.93	C	С
	ATOM	4938		ILE	226		-54.325		1.00 14.79	С	C
	MOTA	4939	С	ILE	226		-54.347		1.00 15.04	С	С
	MOTA	4940	0	ILE	226		-53.970		1.00 14.65	C	0
00	MOTA	4941	N	GLU	227		-55.296		1.00 14.65	C	N
20	ATOM	4942	CA	GLU	227		-55.955		1.00 15.12	С	С
	MOTA	4943	СВ	GLU	227		-57.088		1.00 15.21	С	С
	MOTA	4944	CG	GLU	227		-58.390		1.00 17.13	С	С
	MOTA	4945	CD	GLU	227		-58.216		1.00 18.21	С	С
~=	MOTA	4946		GLU	227		-57.530		1.00 20.90	С	0
25	MOTA	4947		GLU	227			154.804	1.00 18.86	С	0
	MOTA	4948	С	GLU	227		-54.965		1.00 14.83	C	С
	MOTA	4949	0	GLU	227		-55.178		1.00 14.73	С	0
	ATOM '	4950	N	ASP	228			158.717	1.00 14.22	С	N
00	MOTA	4951	CA	ASP	228			159.793	1.00 15.26	C	C
30	MOTA	4952	CB	ASP	228			159.807	1.00 14.81	С	С
	ATOM	4953	CG	ASP	228			160.109	1.00 16.58	С	С
	MOTA	4954		ASP	228			161.069	1.00 15.48	С	0
	MOTA	4955		ASP	228			159.391	1.00 15.15	C	0
0=	ATOM	4956	С	ASP	228			159.636	1.00 15.16	C	С
35	ATOM	4957	0	ASP	228			160.609	1.00 15.37	С	0
	ATOM	4958	N	GLY	229			158.400	1.00 15.16	С	N
	MOTA	4959	CA	GLY	229			158.151	1.00 14.56	С	С
	MOTA	4960	С	GLY	229			158.416	1.00 14.33	C	C
40	MOTA	4961	0	GLY	229			158.987	1.00 12.80	С	0
40	MOTA	4962	N	ALA	230			158.023	1.00 13.05	С	N
	MOTA	4963	CA	ALA	230			158.223	1.00 13.45	С	С
	ATOM	4964	СВ	ALA	230			157.452	1.00 12.06	_	_
	ATOM	4965	С	ALA	230			159.701	1.00 13.54	C	C
45	MOTA	4966	0	ALA	230			160.131	1.00 12.94	С	0
45	MOTA	4967	N	ARG	231			160.482	1.00 13.80	С	N
	MOTA	4968	CA	ARG	231			161.903	1.00 14.40	С	С
	MOTA	4969	СВ	ARG	231			162.536	1.00 15.07	С	С
	MOTA	4970	CG	ARG	231			162.018	1.00 16.54	С	С
50	MOTA	4971	CD	ARG	231			162.238	1.00 16.33	С	C
50	ATOM	4972	NE	ARG	231			163.632	1.00 16.45	С	N
	MOTA	4973	CZ	ARG	231			164.250	1.00 17.78	С	С
	MOTA	4974		ARG	231			163.611	1.00 17.87	С	N
	ATOM	4975		ARG	231			165.514	1.00 18.04	С	N
EE	ATOM	4976	C	ARG	231			162.710	1.00 14.96	С	С
55	ATOM	4977	0	ARG	231			163.797	1.00 14.90	С	0
	MOTA	4978	N	VAL	232			162.207	1.00 15.16	C	N
	MOTA	4979	CA	VAL	232			162.949	1.00 15.82	С	С
	MOTA	4980	CB	VAL	232	-0.277	-50.372	162.862	1.00 15.90	C	C

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	MOTA	4981	CG1	VAL	232	1.126	-50.538	163.444	1.00 15.46	С	С
	MOTA	4982	CG2	VAL	232	-0.222	-49.852	161.421	1.00 13.24	С	C
	MOTA	4983	С	VAL	232	-2.496	-51.562	162.503	1.00 15.62	С	С
	MOTA	4984	0	VAL	232	-3.213	-50.718	163.033	1.00 14.53	С	0
5	ATOM	4985	N	GLY	233	-2.937	-52.367	161.535	1.00 16.24	C	N
	ATOM	4986		GLY	233		-52.284		1.00 17.01	c	C
	ATOM	4987	C	GLY	233		-51.859		1.00 17.97	Ċ	Ċ
	ATOM	4988	Ö	GLY	233		-51.926		1.00 18.13	č	ō
	ATOM	4989	N	PHE	234		-51.411		1.00 10.13	C	N
10	ATOM	4990	CA	PHE	234		-51.016		1.00 17.03		
10		4991			234				· ·	C	C
	MOTA		CB	PHE			-50.295		1.00 17.03	C	C
	MOTA	4992	CG	PHE	234		-48.908	157.421	1.00 18.50	C	C
	MOTA	4993	CD1		234		-48.605		1.00 17.96	C	С
4-	MOTA	4994	CD2		234			157.137	1.00 17.72	C	С
15	MOTA	4995	CE1		234			158.684	1.00 19.80	C	C
	MOTA	4996	CE2	PHE	234			157.616	1.00 18.38	С	С
	MOTA	4997	CZ	PHE	234	-2.207	-46.306	158.393	1.00 20.51	C	С
	MOTA	4998	С	PHE	234	-4.370	-52.235	156.703	1.00 18.19	C	C
	MOTA	4999	0	PHE	234	-3.834	-53.328	156.909	1.00 17.65	С	0
20	MOTA	5000	N	GLN	235	-5.298	-52.046	155.767	1.00 18.33	С	N
	MOTA	5001	CA	GLN	235			154.888	1.00 19.22	C	С
	MOTA	5002	СВ	GLN	235			154.184	1.00 19.75	C	C
	ATOM	5003	CG	GLN	235			155.144	1.00 21.43	Č	Ċ
	ATOM	5004	CD	GLN	235			154.432	1.00 22.69	Č	c
25	ATOM	5005		GLN	235			153.608	1.00 22.47	C	Ö
20	MOTA	5006	NE2		235			154.751	1.00 22.47	C	N
		5007	C	GLN	235						
	MOTA							153.864	1.00 19.00	C	С
	ATOM	5008	0	GLN	235			153.393	1.00 17.75	C	0
20	MOTA	5009	N	VAL	236			153.515	1.00 19.65	С	N
30	MOTA	5010	CA	VAL	236			152.564	1.00 19.44	C	C
	MOTA	5011	CB	VAL	236			152.354	1.00 19.19	С	C
	MOTA	5012		VAL	236			151.320	1.00 17.93	С	С
	MOTA	5013		VAL	236			153.681	1.00 18.12	С	C
0_	MOTA	5014	С	VAL	236	-3.561	-54.360	151.218	1.00 19.53	С	C
35	MOTA	5015	0	VAL	236	-2.554	-53.943	150.648	1.00 18.39	С	0
	MOTA	5016	N	GLU	237	-4.784	-54.251	150.708	1.00 19.52	C	N
	MOTA	5017	ÇA	GLU	237	-5.021	-53.598	149.418	1.00 20.17	С	С
	ATOM	5018	CB	GLU	237	-6.512	-53.631	149.088	1.00 22.85	С	C
	MOTA	5019	CG	GLU	237		-53.098		1.00 27.72	C	C
40	ATOM	5020	CD	GLU	237			147.431	1.00 31.75	C	C
	ATOM	5021	OE1		237			148.270	1.00 33.80	Č	ō
	ATOM	5022		GLU	237			146.371	1.00 33.68	Č	ō
	ATOM	5023	C	GLU	237			149.454	1.00 18.07	C	Č
	ATOM	5024	Ö	GLU	237			148.534	1.00 16.31	Č	ō
45		5025	N	PHE	238			150.515	1.00 16.98		
70	ATOM	5025								C	N
	ATOM		CA	PHE	238			150.693	1.00 17.27	C	C
	ATOM	5027	CB	PHE	238			152.027	1.00 17.34	C	C
	MOTA	5028	CG	PHE	238			152.397	1.00 17.96	C	C
	MOTA	5029		PHE	238			151.694	1.00 18.18	C	С
50	MOTA	5030		PHE	238			153.421	1.00 18.28	С	С
	MOTA	5031		PHE	238			152.006	1.00 17.80	С	C
	MOTA	5032		PHE	238			153.743	1.00 18.89	С	C
	MOTA	5033	CZ	PHE	238			153.032	1.00 18.33	С	CCC
	MOTA	5034	С	PHE	238	-2.863	-50.058	150.703	1.00 16.70	С	С
55	MOTA	5035	0	PHE	238	-2.215	-49.278	150.008	1.00 16.30	С	0
	MOTA	5036	N	LEU	239			151.505	1.00 16.06	С	N
	ATOM	5037	CA	LEU	239			151.615	1.00 17.40	С	C
	MOTA	5038	СВ	LEU	239			152.598	1.00 15.80	Č	Č
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	MOTA	5039		LEU	239	-0.056	-51.903	154.046	1.00 17.46	C	С
	MOTA	5040	CD1		239			154.480	1.00 15.39	C	С
	MOTA	5041	CD2		239		-52.946		1.00 16.59	С	С
_	MOTA	5042	С	LEU	239		-51.335		1.00 17.56	C	C
5	MOTA	5043	0	LEU	239		-50.696		1.00 17.19	C	0
	ATOM	5044	N	GLU	240			149.468	1.00 18.40	С	N
	MOTA	5045	CA	GLU	240			148.145	1.00 20.85	C	C
	MOTA	5046	CB	GLU	240			147.521	1.00 23.02	C	С
	MOTA	5047	CG	GLU	240			148.259	1.00 26.39	С	С
10	MOTA	5048	CD	GLU	240			147.972	1.00 30.97	C	С
	MOTA	5049	OE1		240			147.772	1.00 32.51	C	0
	MOTA	5050	OE2		240			147.958	1.00 33.15	С	0
	MOTA	5051	С	GLU	240			147.230	1.00 20.90	С	С
	MOTA	5052	0	GLU	240			146.454	1.00 20.74	C	0
15	MOTA	5053	N	LEU	241			147.316	1.00 21.07	С	N
	MOTA	5054	CA	LEU	241			146.501	1.00 22.58	C	С
	MOTA	5055	CB	LEU	241			146.741	1.00 24.26	С	C
	MOTA	5056	CG	LEU	241			146.093	1.00 26.53	C	C
	MOTA	5057		LEU	241			144.575	1.00 27.75	C	С
20	MOTA	5058		LEU	241			146.513	1.00 28.98	С	С
	MOTA	5059	С	LEU	241			146.865	1.00 22.18	С	С
	MOTA	5060	0	LEU	241			145.997	1.00 22.71	C	0
	ATOM	5061	N	LEU	242	-0.088	-48.301	148.158	1.00 20.92	C	N
	MOTA	5062	CA	LEU	242			148.628	1.00 20.02	C	С
25	MOTA	5063	CB	LEU	242	0.860	-47.274	150.153	1.00 20.07	С	С
	MOTA	5064	CG	LEU	242			150.806	1.00 21.59	С	C
	MOTA	5065		LEU	242	1.568	-44.864	150.189	1.00 20.97	C	С
	MOTA	5066		LEU	242			152.299	1.00 19.92	C	С
	MOTA	5067	С	LEU	242			148.196	1.00 19.08	С	C
30	MOTA	5068	0	LEU	242			147.680	1.00 18.08	C	0
	MOTA	5069	N	PHE	243			148.396	1.00 18.57	C	N
	ATOM .	5070	CA	PHE	243			148.006	1.00 18.80	С	C
	MOTA	5071	CB	PHE	243			148.603	1.00 16.93	С	C
٥-	MOTA	5072	CG	PHE	243			150.053	1.00 15.95	C	C
35	MOTA	5073	CD1		243			151.049	1.00 15.49	C	C
	MOTA	5074	CD2		243			150.410	1.00 14.54	С	C
	MOTA	5075	CE1		243			152.384	1.00 15.83	C	С
	MOTA	5076	CE2		243			151.738	1.00 15.46	С	С
40	MOTA	5077	CZ	PHE	243			152.732	1.00 14.97	С	С
40	ATOM	5078	С	PHE	243			146.484	1.00 19.80	С	C
	ATOM	5079	0	PHE	243			146.010	1.00 18.39	C	0
	MOTA	5080	N	HIS	244			145.720	1.00 20.82	С	N
	MOTA	5081	CA	HIS	244			144.258	1.00 22.43	С	C
45	ATOM	5082	CB	HIS	244			143.609	1.00 25.14	С	C
45	ATOM	5083	CG	HIS	244			142.112	1.00 29.07	С	C
	ATOM	5084		HIS	244			141.269	1.00 30.07	С	С
	MOTA	5085		HIS	244			141.312	1.00 29.94	С	N
	MOTA	5086		HIS	244			140.042	1.00 30.15	С	С
<b>E</b> 0	MOTA	5087		HIS	244			139.989	1.00 30.65	С	N
50	ATOM	5088	C	HIS	244			143.834	1.00 21.93	С	С
	ATOM	5089	0	HIS	244			142.914	1.00 21.81	С	0
	ATOM	5090	N	PHE	245			144.510	1.00 20.16	С	N
	ATOM	5091	CA	PHE	245			144.240	1.00 19.29	C	С
EE	MOTA	5092	CB	PHE	245			145.195	1.00 19.02	С	C
55	ATOM	5093	CG	PHE	245			145.238	1.00 18.10	C	C
	ATOM	5094		PHE	245			144.242	1.00 18.27	C	C
	ATOM	5095		PHE	245			146.263	1.00 18.82	С	С
	MOTA	5096	CE1	PHE	245	2.928	-41.451	144.264	1.00 18.07	C	C

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	MOTA	5097	CE2	PHE	245	4.142	-41.913	146.300	1.00 18.71	С	С
	MOTA	5098	CZ	PHE	245			145.295	1.00 19.85	C	C
	MOTA	5099	С	PHE	245			144.470	1.00 18.96	C	C
	MOTA	5100	0	PHE	245	5.355	-45.156	143.621	1.00 17.49	Ċ	0
5	ATOM	5101	N	HIS	246			145.633	1.00 17.92	Č	N
	ATOM	5102	CA	HIS	246	6.577	-45.917	145.948	1.00 18.17	C	C
	ATOM	5103	СВ	HIS	246	6.872	-46.353	147.382	1.00 17.39	c	C
	MOTA	5104	CG	HIS	246	6.504	-45.318	148.402	1.00 16.79	Č	C
	MOTA	5105	CD2	HIS	246			149.279	1.00 16.89	C	С
10	ATOM	5106	ND1	HIS	246	7.211	-44.145	148.555	1.00 14.73	С	N
	MOTA	5107	CE1	HIS	246	6.631	-43.400	149.480	1.00 16.89	C	С
	MOTA	5108	NE2	HIS	246			149.936	1.00 16.76	C	N
	ATOM	5109	С	HIS	246			144.962	1.00 17.97	C	С
	ATOM	5110	0	HIS	246	8.511	-45.968	144.560	1.00 17.61	С	0
15	MOTA	5111	N	GLY	247	7.243	-47.803	144.552	1.00 18.68	C	N
	MOTA	5112	CA	GLY	247			143.587	1.00 19.53	C	C
	MOTA	5113	С	GLY	247			142.243	1.00 20.19	Ċ	C
	ATOM	5114	0	GLY	247			141.646	1.00 20.80	Ċ	Ō
	ATOM	5115	N	THR	248	6.921	-47.373	141.765	1.00 19.87	C	N
20	ATOM	5116	CA	THR	248			140.491	1.00 19.32	Ċ	C
	ATOM	5117	СВ	THR	248			140.160	1.00 19.66	Č	Ċ
	ATOM	5118		THR	248			140.176	1.00 19.35	č	ō
	ATOM	5119	CG2		248			138.783	1.00 18.52	Ċ	Ċ
	ATOM	5120	С	THR	248			140.491	1.00 19.29	Č	Ċ
25	ATOM	5121	0	THR	248	8.259	-45.029	139.554	1.00 18.25	Ċ	Ō
	ATOM	5122	N	LEU	249			141.531	1.00 18.21	Č	N
	ATOM	5123	CA	LEU	249			141.604	1.00 19.06	Č	C
	ATOM	5124	CB	LEU	249			142.826	1.00 18.37	Č	Č
	ATOM	5125	CG	LEU	249			143.067	1.00 19.31	Č	Č
30	ATOM	5126		LEU	249			141.886	1.00 17.37	Č	Č
	ATOM	5127		LEU	249			144.356	1.00 19.95	Č	č
	ATOM	5128	С	LEU	249			141.694	1.00 19.08	Ċ	Ċ
	ATOM	5129	Ó	LEU	249			140.998	1.00 18.35	Č	ō
	ATOM	5130	N	ARG	250			142.555	1.00 19.05	C	N
35	ATOM	5131	CA	ARG	250			142.748	1.00 21.51	Ċ	C
	ATOM	5132	CB	ARG	250			143.793	1.00 22.24	C	C
	ATOM	5133	CG	ARG	250			144.286	1.00 25.15	Ċ	Č
	ATOM	5134	CD	ARG	250			145.095	1.00 27.57	Č	c
	ATOM	5135	NE	ARG	250			146.097	1.00 29.42	Č	N
40	ATOM	5136	CZ	ARG	250			145.909	1.00 31.82	c	C
	ATOM	5137	NH1		250			144.747	1.00 33.24	Č	N
	ATOM	5138		ARG	250			146.884	1.00 32.37	Č	N
	MOTA	5139	С	ARG	250			141.448	1.00 22.35	C	C
	MOTA	5140	0	ARG	250			141.182	1.00 21.30	C	Ō
45	ATOM	5141	N	LYS	251			140.640	1.00 23.28	C	N
	ATOM	5142	CA	LYS	251			139.385	1.00 24.95	Č	C
	ATOM	5143	СВ	LYS	251			138.750	1.00 25.59	C	c
	ATOM	5144	CG	LYS	251			139.454	1.00 27.12	Č	Ċ
	ATOM	5145	CD	LYS	251			138.759	1.00 29.13	C	C
50	ATOM	5146	CE	LYS	251			139.620	1.00 30.83	C	C
	ATOM	5147	NZ	LYS	251			139.064	1.00 31.97	C	N
	MOTA	5148	C	LYS	251			138.359	1.00 25.23	Č	C
	ATOM	5149	O	LYS	251			137.439	1.00 25.25	c	Ö
	ATOM	5150	N	LEU	252			138.511	1.00 24.72	Č	N
55	ATOM	5151	CA	LEU	252			137.577	1.00 25.19	c	C
	ATOM	5152	СВ	LEU	252			137.668	1.00 24.25	c	c
	MOTA	5153	CG	LEU	252			137.278	1.00 25.10	C	c
	MOTA	5154		LEU	252			137.611	1.00 24.20	c	c
		_								_	_

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	ATOM	5155	CD2		252		-42.861		1.00 24.17	C	С
	MOTA	5156	С	LEU	252		-42.232		1.00 26.15	С	С
	MOTA	5157	0	LEU	252	13.537	-41.354	137.013	1.00 26.54	С	0
	MOTA	5158	N	GLN	253	13.894	-42.591	138.852	1.00 27.14	C	N
5	MOTA	5159	CA	GLN	253	15.182	-41.979	139.147	1.00 28.03	C	С
	ATOM	5160	CB	GLN	253		-42.492		1.00 29.58	č	Ċ
	ATOM	5161	CG	GLN	253		-43.989		1.00 31.02	c	c
	ATOM	5162	CD	GLN	253		-44.495		1.00 31.02		c
	ATOM	5163		GLN	253		-45.643			C	
10	ATOM	5164		GLN			-43.644		1.00 35.27	C	0
10	ATOM	5165			253				1.00 33.37	C	N
			C	GLN	253		-40.461		1.00 27.47	C	C
	MOTA	5166	0	GLN	253		-39.836		1.00 27.66	С	0
	MOTA	5167	N	LEU	254		-39.870		1.00 27.11	С	N
45	MOTA	5168	CA	LEU	254		-38.428		1.00 26.74	C	С
15	MOTA	5169	CB	LEU	254		-38.018		1.00 25.22	C	С
	MOTA	5170	CG	LEU	254		-38.440		1.00 24.23	С	C
	MOTA	5171	CD1	LEU	254	10.345	-37.659	140.762	1.00 22.94	С	С
	ATOM	5172	CD2	LEU	254	11.402	-38.170	138.557	1.00 22.58	С	С
	MOTA	5173	С	LEU	254	15.278	-37.734	140.528	1.00 27.44	С	С
20	MOTA	5174	0	LEU	254			141.395	1.00 26.90	Ċ	Ō
	ATOM	5175	N	GLN	255		-36.519		1.00 28.34	Ċ	N
	ATOM	5176	CA	GLN	255			140.625	1.00 30.41	č	c
	ATOM	5177	СВ	GLN	255			139.542	1.00 30.41	c	C
	ATOM	5178	CG	GLN	255		-35.532		1.00 32.70		
25	ATOM	5179	CD	GLN	255			137.255		C	C
20	MOTA	5179	OE1		255			137.255	1.00 37.99	C	C
		5180							1.00 39.69	C	0
	MOTA		NE2		255			136.111	1.00 39.40	C	N
	MOTA	5182	C	GLN	255			141.726	1.00 30.12	С	С
20	MOTA	5183	0	GLN	255			141.732	1.00 29.99	С	0
30	MOTA	5184	N	GLU	256			142.658	1.00 30.10	C	N
	MOTA	5185	CA	GLU	256			143.757	1.00 30.45	C	C
	ATOM	5186	CB	GLU	256			144.646	1.00 32.12	C	C
	ATOM	5187	CG	GLU	256	16.720	-32.478	145.946	1.00 35.72	C	C
	ATOM	5188	CD	GLU	256		-32.173		1.00 37.90	C	С
35	ATOM	5189	OE1	GLU	256	18.624	-31.249	146.676	1.00 39.48	С	0
	MOTA	5190	OE2	GLU	256	17.881	-32.867	147.973	1.00 39.28	C	0
	MOTA	5191	С	GLU	256			143.354	1.00 29.53	C	C
	ATOM	5192	0	GLU	256			143.946	1.00 28.88	Ċ	ŏ
	MOTA	5193	N	PRO	257			142.349	1.00 28.61	č	N
40	MOTA	5194	CD	PRO	257			141.589	1.00 28.20	c	C
	ATOM	5195	CA	PRO	257			141.976	1.00 27.69	C	c
	ATOM	5196	СВ	PRO	257			140.820	1.00 27.69	C	
	ATOM	5197	CG	PRO	257			141.169		_	_
	ATOM	5198	C	PRO	257			141.109	1.00 28.71	C	C
45		5199							1.00 26.87	C	C
43	ATOM		0	PRO	257			141.891	1.00 26.59	С	0
	ATOM	5200	N	GLU	258			140.858	1.00 25.85	С	N
	ATOM	5201	CA	GLU	258			140.418	1.00 24.88	C	С
	MOTA	5202	СВ	GLU	258			139.439	1.00 24.17	С	C
	MOTA	5203	CG	GLU	258			138.205	1.00 25.10	C	С
50	MOTA	5204	CD	GLU	258			137.360	1.00 26.23	C	С
	ATOM	5205		GLU	258			137.933	1.00 25.87	С	0
	MOTA	5206	OE2	GLU	258			136.118	1.00 27.53	С	0
	ATOM	5207	С	GLU	258			141.629	1.00 23.84	Ċ	Č
	MOTA	5208	0	GLU	258			141.678	1.00 22.84	c	ō
55	ATOM	5209	N	TYR	259			142.611	1.00 22.35	Č	N
	ATOM	5210	CA	TYR	259			143.827	1.00 22.61	C	C
	ATOM	5211	СВ	TYR	259			144.760	1.00 22.01	C	
	ATOM	5212	CG	TYR	259			144.760			С
	VI OU	3212	CG	TIK	233	12.236	-20.288	T44.9T\	1.00 21.14	С	С

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	MOTA	5213	CD1		259		-37.395		1.00 20.56	С	С
	MOTA	5214	CE1	TYR	259		-38.794		1.00 18.62	C	С
	MOTA	5215	CD2	TYR	259	13.481	-37.217	144.394	1.00 20.34	C	С
_	MOTA	5216	CE2	TYR	259		-38.607		1.00 19.30	C	С
5	MOTA	5217	CZ	TYR	259		-39.389		1.00 19.88	C	С
	MOTA	5218	OH	TYR	259	12.564	-40.761	144.427	1.00 17.21	C	0
	ATOM	5219	С	TYR	259	10.523	-33.217	144.582	1.00 22.27	C	С
	MOTA	5220	0	TYR	259		-33.298		1.00 21.25	С	0
	MOTA	5221	N	VAL	260	11.267	-32.132	144.805	1.00 21.69	C	N
10	ATOM	5222	CA	VAL	260		-31.007		1.00 21.85	C	С
	MOTA	5223	СВ	VAL	260		-29.997		1.00 21.84	C	С
	ATOM	5224		VAL	260	12.886	-30.741	146.804	1.00 23.44	С	С
	MOTA	5225	CG2	VAL	260	12.400	-29.245	144.853	1.00 21.89	С	С
	ATOM	5226	С	VAL	260		-30.256		1.00 21.44	С	С
15	ATOM	5227	0	VAL	260		-29.681		1.00 19.91	С	0
	MOTA	5228	N	LEU	261	9.758	-30.257	143.456	1.00 21.26	С	N
	MOTA	5229	CA	LEU	261	8.782	-29.579	142.609	1.00 23.04	С	С
	MOTA	5230	CB	LEU	261	9.309	-29.508	141.172	1.00 24.60	С	C
	MOTA	5231	CG	LEU	261	9.510	-28.172	140.440	1.00 24.84	С	С
20	MOTA	5232	CD1	LEU	261	9.586	-27.008	141.388	1.00 24.16	С	С
	MOTA	5233	CD2	LEU	261	10.784	-28.282	139.599	1.00 25.17	С	С
	MOTA	5234	С	LEU	261	7.481	-30.377	142.667	1.00 22.74	C	С
	MOTA	5235	0	LEU	261		-29.818		1.00 22.54	C	ō
	ATOM	5236	N	LEU	262		-31.694		1.00 23.07	Ċ	N
25	MOTA	5237	CA	LEU	262		-32.587		1.00 23.17	Č	С
	MOTA	5238	CB	LEU	262	6.950	-34.033	142.863	1.00 24.16	Ċ	Ċ
	ATOM	5239	CG	LEU	262			142.507	1.00 27.12	Č	Č
	ATOM	5240	CD1	LEU	262			141.135	1.00 26.19	Č	Č
	MOTA	5241	CD2	LEU	262			142.514	1.00 26.28	Ċ	Č
30	ATOM	5242	С	LEU	262			144.205	1.00 22.78	Ċ	C
	ATOM	5243	0	LEU	262			144.249	1.00 21.96	č	ŏ
	ATOM	5244	N	ALA	263			145.279	1.00 21.56	Č	N
	ATOM	5245	CA	ALA	263			146.583	1.00 21.33	Ç	C
	ATOM	5246	СВ	ALA	263			147.648	1.00 19.74	Č	Č
35	ATOM	5247	С	ALA	263			146.531	1.00 20.96	c	Č
	MOTA	5248	0	ALA	263			147.148	1.00 20.00	Ċ	ŏ
	ATOM	5249	N	ALA	264			145.798	1.00 20.08	Ċ	N
	ATOM	5250	CA	ALA	264			145.663	1.00 21.05	Č	C
	MOTA	5251	СВ	ALA	264			144.853	1.00 19.86	Č	č
40	MOTA	5252	С	ALA	264	3.943		144.980	1.00 21.31	c	Č
	ATOM	5253	0	ALA	264			145.394	1.00 21.71	Č	ō
	ATOM	5254	N	MET	265			143.928	1.00 20.89	c	
	ATOM	5255	CA	MET	265	2.597	-29.231	143.210	1.00 21.67	Ċ	C
	ATOM	5256	CB	MET	265			141.980	1.00 22.87	c	c
45	ATOM	5257	CG	MET	265			140.838	1.00 25.80	Č	Č
	ATOM	5258	SD	MET	265			139.427	1.00 29.35	Č	S
	ATOM	5259	CE	MET	265			138.513	1.00 26.54	c	C
	MOTA	5260	C	MET	265			144.128	1.00 20.81	C	c
	ATOM	5261	Ö	MET	265			144.066	1.00 19.86	c	0
50	ATOM	5262	N	ALA	266			144.992	1.00 20.98	C	N
••	ATOM	5263	CA	ALA	266			145.922	1.00 21.57	C	
	ATOM	5264	СВ	ALA	266			146.655	1.00 20.93	C	C
	MOTA	5265	C	ALA	266			146.931	1.00 20.93	C	C
	MOTA	5266	Ö	ALA	266			140.931	1.00 21.45	C	0
55	ATOM	5267	N	LEU	267			147.234	1.00 20.27	C	
55	ATOM	5268	CA	LEU	267			148.391	1.00 21.93		N
	ATOM	5269	CB	LEU	267			148.751		C	C
	ATOM	5270	CG	LEU	267			150.145	1.00 23.06	C	C
	AION	3270	CG	LEO	207	د	-20.330	120,142	1.00 23.78	С	C

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	ATOM	5271	CD1	LEU	267	3.619	-25.911	150.100	1.00 2	3.35	С	С
	ATOM	5272	CD2	LEU	267		-26.417		1.00 2		Č	Ċ
	ATOM	5273	C	LEU	267		-27.390		1.00 2		Č	Č
	ATOM	5274	Ö	LEU	267		-27.090		1.00 2		C	ō
5	ATOM	5275	N	PHE	268		-26.870		1.00 2		c	N
•	ATOM	5276	CA	PHE	268		-25.881					
	ATOM	5277	CB	PHE					1.00 2		C	C
					268		-24.870		1.00 2		C	C
	ATOM	5278	CG	PHE	268		-24.097		1.00 2		С	С
40	MOTA	5279	CD1		268		-24.256		1.00 2		С	С
10	MOTA	5280		PHE	268		-23.252		1.00 2		C	С
	MOTA	5281		PHE	268		-23.588		1.00 2		С	С
	MOTA	5282		PHE	268		-22.579		1.00 2	7.94	C	С
	MOTA	5283	CZ	PHE	268	3.167	-22.748	147.678	1.00 2	8.60	C	C
	MOTA	5284	C	PHE	268	-1.542	-26.480	145.155	1.00 2	9.29	С	С
15	ATOM	5285	0	PHE	268	-1.512	-26.351	143.935	1.00 2	8.87	C	0
	ATOM	5286	N	SER	269	-2.495	-27.141	145.800	1.00 3	1.35	С	N
	MOTA	5287	CA	SER	269	-3.625	-27.745	145.115	1.00 3	3.69	С	C
	ATOM	5288	CB	SER	269	-3.858	-29.168	145.616	1.00 3	3.87	С	C
	MOTA	5289	OG	SER	269	-2.838	-30.031	145.148	1.00 3		Ċ	o
20	MOTA	5290	С	SER	269		-26.864		1.00 3		Č	Ċ
	ATOM	5291	0	SER	269		-26.801		1.00 3		č	ō
	ATOM	5292	N	PRO	270		-26.163		1.00 3		c	N
	ATOM	5293	CD	PRO	270		-26.297		1.00 3		c	C
	ATOM	5294	CA	PRO	270		-25.254		1.00 3		c	C
25	ATOM	5295	CB	PRO	270		-24.626			–		
20	ATOM	5296	CG	PRO	270		-24.020		1.00 3		C	C
	ATOM	5297	C						1.00 3		C	C
			-	PRO	270		-25.925		1.00 3		C	C
	ATOM	5298	0	PRO	270			145.514	1.00 3		C	0
20	ATOM	5299	N	ASP	271			144.565	1.00 4		С	N
30	ATOM	5300	CA	ASP	271			144.861	1.00 4		С	C
	ATOM	5301	СВ	ASP	271			143.749	1.004		С	С
	ATOM	5302	CG	ASP	271			143.658	1.004		С	С
	MOTA	5303		ASP	271			143.728	1.004		С	0
0.5	ATOM	5304		ASP	271			143.499	1.00 4	4.16	C	0
35	MOTA	5305	С	ASP	271			146.207	1.004	2.07	С	С
	MOTA	5306	0	ASP	271			146.363	1.00 4	2.01	С	0
	MOTA	5307	N	ARG	272			147.178	1.00 4	2.23	C	N
	MOTA	5308	CA	ARG	272			148.516	1.00 4	2.88	C	C
	MOTA	5309	CB	ARG	272	-7.007	-28.084	149.139	1.00 4	2.11	C	С
40	ATOM	5310	CG	ARG	272	-6.285	-29.134	149.948	1.00 4	1.65	С	С
	MOTA	5311	CD	ARG	272	-5.106	-29.710	149.179	1.00 4	0.41	С	С
	ATOM	5312	NE	ARG	272	-5.240	-31.145	148.964	1.00 3	9.96	C	N
	ATOM	5313	CZ	ARG	272	-4.260	-31.951	148.563	1.00 3	8.64	C	C
	MOTA	5314	NH1	ARG	272	-3.045	-31.480	148.327	1.00 3		C	N
45	ATOM	5315		ARG	272	-4.502	-33.240	148.387	1.00 3		C	N
	ATOM	5316	С	ARG	272			149.385	1.00 4		C	C
	ATOM	5317	0	ARG	272			149.256	1.00 4		Ċ	ō
	ATOM	5318	N	PRO	273			150.284	1.00 4		Č	N
	MOTA	5319	CD	PRO	273			150.507	1.00 4		Č	C
50	MOTA	5320	CA	PRO	273			151.150	1.00 4		c	C
00	MOTA	5321	CB	PRO	273			151.894	1.00 4		c	C
	ATOM	5322	CG	PRO	273			151.894				C
									1.00 4		C	С
	ATOM	5323	C	PRO	273			152.096	1.00 4		C	С
55	MOTA	5324	0	PRO	273			152.898	1.00 4		C	0
55	ATOM	5325	N	GLY	274			151.986	1.00 4		C	N
	ATOM	5326	CA	GLY	274			152.839	1.00 4		С	С
	ATOM	5327	C	GLY	274			152.196	1.00 4		С	С
	MOTA	5328	0	GLY	274	-10.379	-22.908	152.830	1.00 4	18.57	С	0

	MOTA	5329	N	VAL	275	-10.113	-24.095	150.945	1.00 48.85	С	N
	ATOM	5330	CA	VAL	275	-9.411	-23.016	150.252	1.00 49.89	С	C
	MOTA	5331	CB	VAL	275		-23.500		1.00 49.99	С	C
_	MOTA	5332	CG1		275	-7.739	-24.549	149.183	1.00 50.05	С	С
5	MOTA	5333	CG2	VAL	275	-9.912	-24.060	148.025	1.00 50.42	С	С
	MOTA	5334	С	VAL	275	-10.289	-21.804	149.958	1.00 50.42	С	С
	MOTA	5335	0	VAL	275	-11.468	-21.939	149.631	1.00 49.97	C	0
	ATOM	5336	N	THR	276	-9.692	-20.620	150.072	1.00 51.16	С	N
	MOTA	5337	CA	THR	276	-10.389	-19.365	149.813	1.00 51.63	С	С
10	MOTA	5338	СВ	THR	276	-10.118	-18.333	150.928	1.00 51.76	C	C
	MOTA	5339	OG1	THR	276	-10.588	-18.844	152.180	1.00 51.76	С	0
	MOTA	5340	CG2	THR	276	-10.830	-17.027	150.629	1.00 52.34	C	С
	MOTA	5341	С	THR	276	-9.924	-18.788	148.479	1.00 52.17	С	C
	MOTA	5342	0	THR	276	-10.722	-18.611	147.558	1.00 52.52	С	0
15	ATOM	5343	N	GLN	277	-8.629	-18.498	148.378	1.00 52.57	С	N
	MOTA	5344	CA	GLN	277	-8.053	-17.951	147.154	1.00 52.67	С	С
	MOTA	5345	CB	GLN	277		-17.195		1.00 53.56	C	C
	MOTA	5346	CG	GLN	277	-6.913	-15.676	147.596	1.00 55.26	C	С
	ATOM	5347	CD	GLN	277	-7.875	-15.252	148.694	1.00 56.34	C	C
20	ATOM	5348	OE1	GLN	277	-9.070	-15.547	148.641	1.00 56.71	C	0
	ATOM	5349	NE2	GLN	277			149.698	1.00 56.69	C	N
	MOTA	5350	С	GLN	277			146.144	1.00 52.40	C	C
	ATOM	5351	0	GLN	277	-6.619	-19.374	145.847	1.00 51.95	С	0
	MOTA	5352	N	ARG	278	-8.839	-19.654	145.609	1.00 52.21	C	N
25	MOTA	5353	CA	ARG	278	-8.720	-20.740	144.644	1.00 52.47	C	С
	MOTA	5354	CB	ARG	278	-10.111	-21.228	144.226	1.00 54.03	C	C
	ATOM	5355	CG	ARG	278	-10.105	-22.600	143.568	1.00 56.61	C	C
	MOTA	5356	CD	ARG	278	-11.488	-23.005	143.074	1.00 58.62	C	С
	ATOM	5357	NE	ARG	278	-11.550	-24.434	142.770	1.00 60.54	C	N
30	ATOM	5358	CZ	ARG	278	-12.604	-25.056	142.245	1.00 61.18	С	С
	ATOM	5359	NH1	ARG	278	-13.712	-24.384	141.949	1.00 61.12	С	N
	ATOM	5360	NH2	ARG	278	-12.552	-26.364	142.022	1.00 61.81	С	N
	ATOM	5361	С	ARG	278	-7.922	-20.357	143.400	1.00 51.41	С	C
	MOTA	5362	0	ARG	278	-7.084	-21.129	142.933	1.00 51.31	C	0
35	ATOM	5363	N	ASP	279	-8.182	-19.170	142.861	1.00 50.43	C	N
	ATOM	5364	CA	ASP	279			141.668	1.00 48.92	С	C
	MOTA	5365	СВ	ASP	279	-8.122	-17.431	141.123	1.00 50.20	C	С
	ATOM	5366	CG	ASP	279			140.530	1.00 51.42	С	C
	MOTA	5367		ASP	279			141.296	1.00 52.30	С	0
40	ATOM	5368	OD2	ASP	279	-9.640	-17.568	139.294	1.00 52.68	С	0
	MOTA	5369	С	ASP	279			141.916	1.00 47.51	С	C
	MOTA	5370	0	ASP	279				1.00 47.56	С	0
	MOTA	5371	N	GLU	280			143.019	1.00 46.03	C	N
	MOTA	5372	CA	GLU	280			143.339	1.00 44.77	С	C
45	ATOM	5373	CB	GLU	280			144.614	1.00 46.49	С	C
	MOTA	5374	CG	GLU	280	-2.776	-16.603	145.185	1.00 48.88	С	С
	ATOM	5375	CD	GLU	280	-2.682	-15.619	146.331	1.00 50.21	С	С
	MOTA	5376		GLU	280			147.141	1.00 51.50	С	0
	MOTA	5377	OE2	GLU	280			146.430	1.00 51.27	C	0
50	MOTA	5378	С	GLU	280			143.501	1.00 42.73	C	С
	MOTA	5379	0	GLU	280			142.994	1.00 42.02	C	0
	MOTA	5380	N	ILE	281	-4.069	-19.805	144.206	1.00 41.11	C	N
	MOTA	5381	CA	ILE	281			144.414	1.00 40.07	С	C
	ATOM	5382	СВ	ILE	281	-4.193	-21.952	145.439	1.00 38.98	С	С
55	MOTA	5383		ILE	281			145.480	1.00 37.84	С	С
	MOTA	5384		ILE	281			146.820	1.00 38.21	С	С
	MOTA	5385		ILE	281			147.882	1.00 38.30	С	С
	ATOM	5386	С	ILE	281	-3.313	-21.835	143.081	1.00 40.18	С	С

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	MOTA	5387	0	ILE	281	-2.318	-22.508	142.810	1.00 39.57	С	0
	MOTA	5388	N	ASP	282	-4.344	-21.692	142.252	1.00 40.74	C	N
	MOTA	5389	CA	ASP	282	-4.390	-22.319	140.934	1.00 41.64	С	С
	MOTA	5390	СB	ASP	282	-5.695	-21.949	140.226	1.00 44.29	С	С
5	ATOM	5391	CG	ASP	282	-6.024	-22.882	139.077	1.00 46.45	C	С
	ATOM	5392	OD1	ASP	282	-5.120	-23.187	138.270	1.00 48.25	С	0
	ATOM	5393	OD2		282		-23.305		1.00 48.22	Č	0
	ATOM	5394	С	ASP	282		-21.835		1.00 41.04	C	Č
	ATOM	5395	Ö	ASP	282		-22.629		1.00 40.33	C	Ö
10	MOTA	5396	N	GLN	283		-20.521				
10		5397							1.00 40.82	C	N
	ATOM		CA	GLN	283			139.358	1.00 40.46	С	С
	MOTA	5398	CB	GLN	283			139.493	1.00 41.88	С	С
	MOTA	5399	CG	GLN	283			139.074	1.00 45.42	С	С
45	MOTA	5400	CD	GLN	283			137.591	1.00 47.59	С	С
15	MOTA	5401		GLN	283			137.062	1.00 48.47	C	0
	MOTA	5402	NE2	GLN	283	-0.279	-16.652	136.913	1.00 49.11	C	N
	MOTA	5403	С	GLN	283	-0.557	-20.408	139.879	1.00 39.02	С	С
	MOTA	5404	0	GLN	283	0.381	-20.626	139.110	1.00 38.65	С	0
	MOTA	5405	N	LEU	284	-0.470	-20.574	141.194	1.00 37.60	C	N
20	ATOM	5406	CA	LEU	284			141.824	1.00 36.30	Ċ	C
	ATOM	5407	СВ	LEU	284			143.346	1.00 36.28	C	Č
	ATOM	5408	CG	LEU	284			144.172	1.00 36.25	C	C
	ATOM	5409		LEU	284			143.366	1.00 35.74	C	C
	ATOM	5410		LEU	284						
25								145.404	1.00 34.66	C	C
25	ATOM	5411	C	LEU	284			141.349	1.00 35.48	С	С
	ATOM	5412	0	LEU	284			141.044	1.00 34.73	С	0
	MOTA	5413	N	GLN	285			141.272	1.00 35.23	С	N
	MOTA	5414	CA	GLN	285			140.828	1.00 35.83	С	С
	MOTA	5415	СВ	GLN	285	-0.968	-25.549	140.923	1.00 36.09	С	C
30	MOTA	5416	CG	GLN	285	-0.661	-27.036	140.807	1.00 37.57	С	C
	MOTA	5417	CD	GLN	285	-1.863	-27.918	141.050	1.00 39.14	С	C
	ATOM	5418	OE1	GLN	285	-1.730	-29.033	141.555	1.00 40.08	С	0
	ATOM	5419	NE2	GLN	285	-3.047	-27.433	140.681	1.00 39.83	С	N
	ATOM	5420	С	GLN	285			139.401	1.00 35.60	C	C
35	ATOM	5421	0	GLN	285			139.096	1.00 35.31	C	ō
	ATOM	5422	N	GLU	286			138.526	1.00 35.27	č	N
	ATOM	5423	CA	GLU	286			137.142	1.00 35.56	C	C
	ATOM	5424	СВ	GLU	286			136.338	1.00 33.50	C	C
	ATOM	5425	CG	GLU	286			134.997	1.00 37.33		
40	ATOM	5426	CD	GLU	286					C	C
70	ATOM	5427						133.838	1.00 44.16	C	C
				GLU	286			133.674	1.00 45.97	С	0
	ATOM	5428		GLU	286			133.078	1.00 45.32	С	_
	ATOM	5429	C	GLU	286			137.110	1.00 34.24	C	С
4 ==	ATOM	5430	0	GLU	286			136.325	1.00 33.97	C	0
45	ATOM	5431	N	GLU	287			137.968	1.00 33.47	C	N
	ATOM	5432	CA	GLU	287			138.048	1.00 33.05	С	C
	MOTA	5433	CB	GLU	287			139.131	1.00 34.71	С	C
	ATOM	5434	CG	GLU	287	3.671	-19.525	138.713	1.00 38.06	C	С
	ATOM	5435	CD	GLU	287	3.826	-18.528	139.858	1.00 40.74	С	C C
50	ATOM	5436	OE1	GLU	287	4.883	-18.556	140.535	1.00 41.79	C	0
	ATOM	5437		GLU	287			140.079	1.00 41.29	Č	ŏ
	ATOM	5438	C	GLU	287			138.389	1.00 31.74	C	c
	ATOM	5439	Ö	GLU	287			137.805	1.00 31.74	c	0
	MOTA	5440	N	MET	288			137.803	1.00 31.68	C	
55	MOTA	5441	CA	MET	288			139.347			N
55	MOTA	5442	CB	MET	288				1.00 29.43	C	C
								140.936	1.00 29.48	C	C
	MOTA	5443	CG	MET	288			142.175	1.00 30.45	C	C
	MOTA	5444	SD	MET	288	5.726	-24.287	142.830	1.00 31.96	C	S

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	MOTA	5445	CE	MET	288		-22.620		1.00 33.18	C	C
	ATOM	5446		MET	288		-26.160		1.00 28.27	С	C
	ATOM	5447		MET	288		-26.608		1.00 27.19	С	0
_	MOTA	5448	N	ALA	289		-26.478		1.00 28.27	С	N
5	MOTA	5449	CA	ALA	289		-27.399		1.00 29.24	C	C
	MOTA	5450	CB	ALA	289	2.585	-27.492	136.369	1.00 28.71	С	С
	ATOM	5451	С	ALA	289	4.944	-26.983	135.719	1.00 29.75	C	С
	ATOM	5452	0	ALA	289	5.729	-27.786	135.216	1.00 29.90	C	0
•	MOTA	5453	N	LEU	290	4.838	-25.725	135.309	1.00 30.94	С	N
10	ATOM	5454	CA	LEU	290	5.662	-25.196	134.233	1.00 32.24	С	С
	ATOM	5455	СВ	LEU	290			134.000	1.00 33.16	C	C
	ATOM	5456	CG	LEU	290			132.686	1.00 35.16	C	C
	MOTA	5457	CD1	LEU	290			132.427	1.00 35.72	C	C
	ATOM	5458	CD2		290			132.746	1.00 36.06	Č.	C
15	MOTA	5459	C	LEU	290			134.569	1.00 32.67	č	C
. •	ATOM	5460	Ö	LEU	290			133.734	1.00 32.07	C	ŏ
	ATOM	5461	N	THR	291			135.800	1.00 32.93	C	N
	ATOM	5462	CA	THR	291			136.234	1.00 32.97		
	MOTA	5463	CB	THR	291					C	C
20			_	THR				137.683	1.00 33.56	C	C
20	ATOM	5464			291			137.783	1.00 32.87	C	0
	ATOM	5465	CG2	THR	291			138.092	1.00 32.59	С	С
	MOTA	5466	C	THR	291			136.147	1.00 33.12	С	C
	MOTA	5467	0	THR	291			135.722	1.00 32.77	C	0
0.5	MOTA	5468	N	LEU	292			136.556	1.00 33.66	C	N
25	ATOM	5469	CA	LEU	292			136.500	1.00 33.98	С	С
	MOTA	5470	CB	LEU	292	7.641	-29.768	137.092	1.00 34.42	C	C
	ATOM	5471	CG	LEU	292			137.659	1.00 34.71	C	С
	ATOM	5472	CD1	LEU	292	6.719	-31.972	137.772	1.00 34.26	C	С
	ATOM	5473	CD2	LEU	292	8.982	-31.871	136.782	1.00 34.52	С	С
30	MOTA	5474	С	LEU	292	8.999	-29.339	135.042	1.00 34.02	С	С
	ATOM	5475	0	LEU	292			134.717	1.00 33.46	C	O
	ATOM	5476	N	GLN	293			134.173	1.00 34.86	Č	N
	ATOM	5477	CA	GLN	293			132.752	1.00 36.35	C	C
	ATOM	5478	CB	GLN	293			131.991	1.00 36.93	č	Č
35	ATOM	5479	CG	GLN	293			132.436	1.00 38.19	Č	Č
	ATOM	5480	CD	GLN	293			131.612	1.00 39.76	c	c
	ATOM	5481		GLN	293			130.409	1.00 39.18	c	ŏ
	ATOM	5482	NE2		293			132.254	1.00 39.83	C	N
	ATOM	5483	C	GLN	293			132.161	1.00 37.04	C	C
40	ATOM	5484	Õ	GLN	293			131.540	1.00 37.04	c	o
40	ATOM	5485	N	SER	294			132.353		C	
	ATOM	5486	CA	SER	294			132.353	1.00 38.14	-	N
		5487	CB		294				1.00 39.42	C	С
	MOTA	5488	OG	SER				132.343	1.00 39.88	C	C
45	ATOM			SER	294			131.897	1.00 42.89	C	0
45	ATOM	5489	C	SER	294			132.333	1.00 39.45	C	C
	ATOM	5490	0	SER	294			131.559	1.00 39.68	С	0
	MOTA	5491	N	TYR	295			133.607	1.00 39.47	С	N
	MOTA	5492	CA	TYR	295			134.160	1.00 39.54	C	C
	MOTA	5493	CB	TYR	295			135.674	1.00 37.59	С	С
50	MOTA	5494	CG	TYR	295			136.310	1.00 35.04	C	C
	MOTA	5495		TYR	295			136.236	1.00 33.15	С	C
	MOTA	5496		TYR	295			136.787	1.00 32.64	С	C
	MOTA	5497	CD2	TYR	295			136.955	1.00 33.46	С	С
	MOTA	5498	CE2	TYR	295	16.370	-29.170	137.506	1.00 31.92	С	С
55	MOTA	5499	CZ	TYR	295			137.420	1.00 32.53	C	C
	MOTA	5500	ОН	TYR	295			137.968	1.00 33.06	Ċ	ō
	MOTA	5501	С	TYR	295	13.445	-29.736	133.501	1.00 40.70	Ċ	Č
	ATOM	5502	Ō	TYR	295			133.283	1.00 41.09	C	Ö
			-		_		10/	203	2.00 31.00	_	9

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	MOTA	5503	N	ILE	296		-30.445		1.00 42.17	C	N
	MOTA	5504	CA	ILE	296		-31.750		1.00 43.74	С	С
	MOTA	5505	CB	ILE	296		-32.538		1.00 42.58	C	С
_	MOTA	5506	CG2		296		-33.760		1.00 41.40	C	С
5	MOTA	5507	CG1	ILE	296		-32.966		1.00 41.48	C	С
	MOTA	5508		ILE	296			134.268	1.00 39.89	С	С
	MOTA	5509	С	ILE	296			131.089	1.00 45.95	С	С
	MOTA	5510	0	ILE	296			130.593	1.00 45.55	C	0
	MOTA	5511	N	LYS	297			130.399	1.00 48.91	C	N
10	MOTA	5512	CA	LYS	297			129.008	1.00 52.23	С	C
	ATOM	5513	СВ	LYS	297			128.441	1.00 51.99	C	С
	MOTA	5514	CG	LYS	297	_		128.141	1.00 52.84	С	С
	MOTA	5515	CD	LYS	297			127.281	1.00 53.70	C	С
	MOTA	5516	CE	LYS	297			126.904	1.00 54.12	С	С
15	MOTA	5517	NZ	LYS	297			128.051	1.00 54.87	С	N
	MOTA	5518	С	LYS	297			129.014	1.00 54.53	C	С
	MOTA	5519	0	LYS	297			128.441	1.00 55.09	C	0
	MOTA	5520	N	GLY	298			129.661	1.00 57.15	C	N
	MOTA	5521	CA	GLY	298			129.748	1.00 60.22	С	С
20	MOTA	5522	С	GLY	298			130.411	1.00 62.54	С	С
	MOTA	5523	0	GLY	298			131.358	1.00 62.69	C	0
	MOTA	5524	N	GLN	299			129.922	1.00 65.17	С	N
	MOTA	5525	CA	GLN	299			130.525	1.00 67.96	С	С
	ATOM	5526	CB	GLN	299			132.033	1.00 68.38	С	C
25	MOTA	5527	CG	GLN	299			132.776	1.00 69.15	С	С
	MOTA	5528	CD	GLN	299			134.270	1.00 69.43	С	С
	MOTA	5529	OE1		299			134.788	1.00 69.78	С	0
	MOTA	5530		GLN	299			134.973	1.00 69.64	С	N
	MOTA	5531	С	GLN	299			130.246	1.00 69.68	С	С
30	MOTA	5532	0	GLN	299			131.167	1.00 69.64	С	0
	MOTA	5533	N	GLN	300			128.961	1.00 71.61	С	N
	ATOM	5534	CA	GLN	300			128.497	1.00 73.35	С	С
	MOTA	5535	CB	GLN	300			127.502	1.00 73.72	С	С
	ATOM	5536	CG	GLN	300			126.322	1.00 74.58	С	С
35	MOTA	5537	CD	GLN	300			125.683	1.00 74.94	С	С
	MOTA	5538	OE1		300			125.518	1.00 74.96	C	0
	ATOM	5539	NE2		300			125.313	1.00 75.16	С	N
	MOTA	5540	С	GLN	300			127.817	1.00 74.46	C	С
40	ATOM	5541	0	GLN	300			126.943	1.00 74.52	С	0
40	MOTA	5542	N	ARG	301		-35.433		1.00 75.88	C	N
	MOTA	5543	CA	ARG	301			127.663	1.00 77.07	C	C
	ATOM	5544	CB	ARG	301			128.128	1.00 77.84	C	C
	ATOM	5545	CG	ARG	301			127.347	1.00 78.90	C	C
45	ATOM	5546	CD	ARG	301			128.082	1.00 79.88	C	С
45	MOTA	5547	NE	ARG	301			129.060	1.00 80.47	C	N
	MOTA	5548	CZ	ARG	301			128.754	1.00 80.79	C	С
	ATOM	5549		ARG	301			127.488	1.00 80.67	C	N
	MOTA	5550	NH2		301			129.723	1.00 81.13	C	N
<b>E</b> 0	ATOM	5551	C	ARG	301			127.991	1.00 77.29	С	C
50	MOTA	5552	0	ARG	301			127.516	1.00 77.30	С	0
	ATOM	5553	N	ARG	302			128.809	1.00 77.38	C	N
	ATOM	5554	CA	ARG	302			3 129.179	1.00 77.29	C	С
	ATOM	5555	CB	ARG	302			2 130.036		C	C
	ATOM	5556	CG	ARG	302			130.578		C	C
55	MOTA	5557	CD	ARG	302			1 131.998		С	C
	MOTA	5558	NE	ARG	302			1 132.042		С	N
	ATOM	5559		ARG	302			131.754		С	С
	MOTA	5560	NH1	. ARG	302	17.931	-35.950	131.406	1.00 79.26	С	N

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	MOTA	5561	NH2	ARG	302	20.148	-36.384	131.810	1.00 79.38	С	N
	MOTA	5562	C	ARG	302		-40.723		1.00 76.92	С	С
	MOTA	5563	0	ARG	302	18.585	-40.166	126.794	1.00 77.08	C	0
	MOTA	5564	N	PRO	303		-41.991		1.00 76.31	С	N
5	MOTA	5565	CD	PRO	303	18.164	-43.054	128.938	1.00 76.37	C	С
	MOTA	5566	CA	PRO	303	17.673	-42.587	126.631	1.00 75.50	С	С
	ATOM	5567	CB	PRO	303	17.382	-44.051	126.964	1.00 75.85	С	С
	ATOM	5568	CG	PRO	303	18.289	-44.303	128.053	1.00 76.19	C	С
	ATOM	5569	С	PRO	303	16.497	-41.892	125.957	1.00 74.45	С	С
10	MOTA	5570	0	PRO	303	16.409	-41.881	124.740	1.00 74.66	C	0
	MOTA	5571	N	ARG	304	15.590	-41.321	126.748	1.00 72.88	C	N
	MOTA	5572	CA	ARG	304	14.445	-40.604	126.185	1.00 71.00	С	С
	ATOM	5573	CB	ARG	304	13.553	-41.580	125.396	1.00 71.95	С	С
	ATOM	5574	CG	ARG	304	13.004	-42.797	126.185	1.00 72.83	C	С
15	MOTA	5575	CD	ARG	304	11.451	-42.760	126.214	1.00 73.75	C	C
	ATOM	5576	NE	ARG	304	10.894	-42.747	124.857	1.00 74.86	С	N
	MOTA	5577	CZ	ARG	304	10.661	-41.650	124.131	1.00 75.42	С	С
	MOTA	5578	NH1	ARG	304	10.905	-40.437	124.613	1.00 75.19	С	N
	MOTA	5579	NH2		304	10.238	-41.763	122.880	1.00 75.41	С	N
20	MOTA	5580	С	ARG	304	13.588	-39.801	127.189	1.00 68.99	C	С
	MOTA	5581	0	ARG	304			127.391	1.00 69.10	С	0
	ATOM	5582	N	ASP	305			127.783	1.00 66.12	С	N
	ATOM	5583	CA	ASP	305			128.758	1.00 62.53	С	С
	ATOM	5584	CB	ASP	305			130.145	1.00 64.03	С	С
25	ATOM	5585	CG	ASP	305			130.972	1.00 65.08	C	С
	ATOM	5586		ASP	305	10.994	-41.568	131.169	1.00 65.82	C	0
	ATOM	5587		ASP	305	13.167	-41.689	131.418	1.00 65.99	C	0
	ATOM	5588	C	ASP	305			128.422	1.00 59.48	C	C
	ATOM	5589	Õ	ASP	305			129.017	1.00 59.18	C	0
30	ATOM	5590	N	ARG	306			127.450	1.00 55.57	С	N
•	ATOM	5591	CA	ARG	306			127.069	1.00 51.44	C	C
	MOTA	5592	СВ	ARG	306			125.568	1.00 53.15	C	С
	ATOM	5593	CG	ARG	306			125.009	1.00 55.42	С	С
	ATOM	5594	CD	ARG	306			123.755	1.00 57.72	C	C
35	ATOM	5595	NE	ARG	306			122.651	1.00 59.57	С	N
••	ATOM	5596	CZ	ARG	306			121.560	1.00 60.42	С	C
	ATOM	5597	_	ARG	306			121.445	1.00 61.04	C	N
	ATOM	5598	NH2		306	9.463	-38.620	120.579	1.00 61.02	С	N
	ATOM	5599	C	ARG	306			127.671	1.00 47.25	C	C
40	ATOM	5600	ō	ARG	306			127.544	1.00 45.71	C	0
	ATOM	5601	Ŋ	PHE	307			128.327	1.00 42.87	С	N
	ATOM	5602	CA	PHE	307			129.004	1.00 38.61	С	С
	ATOM	5603	СВ	PHE	307			128.787	1.00 38.35	C	C
	ATOM	5604	CG	PHE	307			127.363	1.00 38.97	C	C
45	ATOM	5605		PHE	307			126.876	1.00 38.81	Č	Č
	ATOM	5606		PHE	307			126.497	1.00 38.13	Č	C
	MOTA	5607		PHE	307			125.544	1.00 38.93	Č	Ċ
	ATOM	5608		PHE	307			125.165	1.00 38.71	Ċ	c
	MOTA	5609	CZ	PHE	307			124.687	1.00 38.27	C	C
50	MOTA	5610	c	PHE	307			130.499	1.00 35.65	c	Ċ
00	ATOM	5611	ŏ	PHE	307			131.273	1.00 35.24	C	ō
	ATOM	5612		LEU	308			130.893		Č	Ŋ
	ATOM	5613		LEU	308			132.299		c	C
	ATOM	5614		LEU	308			132.459		c	C
55	ATOM	5615		LEU	308			133.749		c	C
-	ATOM	5616		LEU	308			134.243		c	C
	ATOM	5617		LEU	308			L 134.833		c	C
	MOTA	5618		LEU	308			5 132.981			
	V 1 Old	2010	_	U	200	1.224	-21.40.	<i>,</i>	I. JU 20.JE	_	_

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	MOTA	5619	0	LEU	308	6.660	-37.796	134.026	1.00 26.62	С	0
	MOTA	5620	N	TYR	309		-36.285		1.00 26.74	C	N
	MOTA	5621	CA	TYR	309		-35.317		1.00 26.38	С	С
_	MOTA	5622	CB	TYR	309		-34.046		1.00 26.27	С	С
5	MOTA	5623	CG	TYR	309		-32.974		1.00 26.31	С	С
	MOTA	5624		TYR	309		-32.390		1.00 25.78	С	С
	MOTA	5625		TYR	309		-31.424		1.00 26.17	C	С
	MOTA	5626		TYR	309		-32.561		1.00 26.61	С	С
	ATOM	5627	CE2	TYR	309		-31.593		1.00 26.55	С	С
10	MOTA	5628	CZ	TYR	309		-31.033		1.00 26.04	С	С
	MOTA	5629	ОН	TYR	309		-30.088		1.00 26.95	C	0
	MOTA	5630	С	TYR	309		-35.901		1.00 25.65	С	С
	MOTA	5631	0	TYR	309		-35.805		1.00 24.69	С	0
	MOTA	5632	N	ALA	310		-36.507		1.00 25.00	C	N
15	MOTA	5633	CA	ALA	310		-37.113		1.00 25.46	С	C
	MOTA	5634	CB	ALA	310		-37.742		1.00 23.32	С	С
	ATOM	5635	C	ALA	310		-38.167		1.00 25.38	C	С
	MOTA	5636	0	ALA	310			133.919	1.00 24.96	С	0
	MOTA	5637	N	LYS	311	3.823	-38.997	133.341	1.00 25.22	С	N
20	MOTA	5638	CA	LYS	311	3.835	-40.019	134.386	1.00 25.29	С	С
	MOTA	5639	CB	LYS	311	5.072	-40.909	134.264	1.00 27.18	C	С
	MOTA	5640	CG	LYS	311	5.041	-41.883	133.108	1.00 29.62	С	C
	ATOM	5641	CD	LYS	311	6.294	-42.744	133.111	1.00 32.10	С	C
	MOTA	5642	CE	LYS	311	6.291	-43.739	131.963	1.00 33.43	C	C
25	MOTA	5643	NZ	LYS	311	7.545	-44.543	131.967	1.00 36.49	С	N
	MOTA	5644	С	LYS	311	3.817	-39.386	135.785	1.00 24.36	C	С
	ATOM	5645	0	LYS	311	3.206	-39.927	136.709	1.00 23.40	С	0
	ATOM	5646	N	LEU	312	4.491	-38.250	135.938	1.00 22.72	C	N
	ATOM	5647	CA	LEU	312	4.525	-37.566	137.227	1.00 22.36	C	С
30	MOTA	5648	СВ	LEU	312	5.615	-36.487	137.243	1.00 20.77	C	C
	MOTA	5649	CG	LEU	312	7.066	-36.994	137.252	1.00 20.74	С	C
	ATOM	5650	CD1	LEU	312	8.025	-35.805	137.282	1.00 19.34	С	C
	ATOM	5651	CD2	LEU	312	7.300	-37.895	138.474	1.00 20.28	С	C
	MOTA	5652	С	LEU	312	3.160	-36.961	137.568	1.00 22.23	С	C
<b>35</b> .	MOTA	5653	0	LEU	312	2.787	-36.889	138.739	1.00 22.33	С	0
	ATOM	5654	N	LEU	313	2.415	-36.517	136.562	1.00 21.95	C	N
	MOTA	5655	CA	LEU	313	1.087	-35.978	136.835	1.00 22.37	С	С
	ATOM	5656	CB	LEU	313	0.499	-35.292	135.597	1.00 21.79	С	C
	ATOM	5657	CG	LEU	313			135.184	1.00 22.17	С	C
40	ATOM	5658	CD1	LEU	313	0.375	-33.363	134.037	1.00 22.22	С	C
	MOTA	5659	CD2	LEU	313	1.200	-33.010	136.371	1.00 21.19	C	C
	ATOM	5660	С	LEU	313	0.200	-37.146	137.266	1.00 22.14	С	C
	MOTA	5661	0	LEU	313			138.180	1.00 22.77	С	0
	MOTA	5662	N	GLY	314	0.368	-38.288	3 136.609	1.00 21.90	C	N
45	MOTA	5663	CA	GLY	314	-0.409	-39.458	3 136.965	1.00 22.13	C	С
	MOTA	5664	С	GLY	314	-0.117	-39.885	138.393	1.00 22.23	С	С
	MOTA	5665	0	GLY	314	-1.022	-40.283	3 139.124	1.00 21.32	С	0
	MOTA	5666	N	LEU	315	1.153	-39.809	138.790	1.00 22.49	С	N
	MOTA	5667	CA	LEU	315	1.550	-40.183	3 140.140	1.00 22.84	С	С
50	MOTA	5668	CB	LEU	315	3.077	-40.249	140.250	1.00 23.35	С	C
	MOTA	5669	CG	LEU	315			5 139.587	1.00 23.08	С	С
	ATOM	5670	CD	L LEU	315	5.189	-41.400	139.500	1.00 22.58	С	C
	MOTA	5671	CD	LEU	315			5 140.396		С	С
	MOTA	5672	С	LEU	315	0.978	39.208	3 141.158	1.00 23.01	С	С
55	MOTA	5673	0	LEU	315			142.262		С	0
	MOTA	5674		LEU	316	0.904	-37.929	9 140.793	1.00 22.90	C	N
	MOTA	5675	CA	LEU	316			141.686		С	С
	MOTA	5676		LEU	316			3 141.116		С	

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	MOTA	5677		LEU	316	1.871 -34.892 141.360 1.00 23			C
	MOTA	5678	CD1		316	1.971 -33.540 140.669 1.00 23			С
	MOTA	5679	CD2		316	2.097 -34.745 142.861 1.00 23			С
_	MOTA	5680		LEU	316	-1.148 -37.250 141.884 1.00 23			С
5	MOTA	5681	0	LEU	316	-1.690 -37.080 142.981 1.00 22			0
	MOTA	5682	N	ALA	317	-1.797 -37.715 140.817 1.00 23			N
	MOTA	5683	CA	ALA	317	-3.206 -38.089 140.888 1.00 24			C
	MOTA	5684	CB	ALA	317	-3.747 -38.378 139.489 1.00 23			С
4.0	MOTA	5685	C	ALA	317	-3.335 -39.331 141.775 1.00 2			С
10	MOTA	5686	0	ALA	317	-4.238 -39.424 142.606 1.00 2		С	0
	MOTA	5687	N	GLU	318	-2.421 -40.279 141.603 1.00 2			N
	MOTA	5688	CA	GLU	318	-2.438 -41.500 142.398 1.00 2		С	С
	ATOM	5689	СВ	GLU	318	-1.353 -42.469 141.918 1.00 2		C	C
4.5	MOTA	5690	CG	GLU	318	-1.413 -43.829 142.596 1.00 3		С	С
15	MOTA	5691	CD	GLU	318	-0.543 -44.866 141.906 1.00 3		C	С
	MOTA	5692		GLU	318	-0.627 -44.984 140.665 1.00 3		С	0
	MOTA	5693		GLU	318	0.215 -45.571 142.603 1.00 3		С	0
	MOTA	5694	С	GLU	318	-2.227 -41.194 143.877 1.00 2		С	C
	MOTA	5695	0	GLU	318	-2.849 -41.810 144.740 1.00 2		С	0
20	MOTA	5696	N	LEU	319	-1.347 -40.241 144.165 1.00 2		С	N
	MOTA	5697	CA	LEU	319	-1.066 -39.850 145.541 1.00 2		C	C
	MOTA	5698	СВ	LEU	319	0.106 -38.866 145.564 1.00 2		С	С
	ATOM	5699	ĊG	LEU	319	0.728 -38.525 146.917 1.00 2		С	C
<b>~</b> =	ATOM	5700		LEU	319	1.006 -39.791 147.707 1.00 2		С	С
25	MOTA	5701		LEU	319	2.011 -37.730 146.683 1.00 2		С	С
	MOTA	5702	С	LEU	319	-2.330 -39.226 146.144 1.00 2		C	С
	MOTA	5703	0	LEU	319	-2.593 -39.340 147.340 1.00 2		С	0
	MOTA	5704	N	ARG	320	-3.114 -38.569 145.297 1.00 2		С	N
~~	MOTA	5705	CA	ARG	320	-4.364 -37.967 145.727 1.00 2		C	С
30	MOTA	5706	CB	ARG	320	-4.968 -37.168 144.577 1.00 2		C	C
	MOTA	5707	CG	ARG	320	-6.167 -36.349 144.952 1.00 3		С	С
	MOTA	5708	CD	ARG	320	-5.770 -34.977 145.472 1.00 3		С	C
	MOTA	5709	NE	ARG	320	-6.959 -34.240 145.887 1.00 3		С	N
0.5	MOTA	5710	CZ	ARG	320	<b>-7.089 -32.919 145.846 1.00 3</b>		С	C
35	ATOM	5711	NH1		320	-6.096 -32.157 145.403 1.00 3		С	N
	MOTA	5712	NH2		320	-8.222 -32.363 146.254 1.00 3		С	N
	MOTA	5713	С	ARG	320	-5.316 -39.099 146.148 1.00 2		C	С
	ATOM	5714	0	ARG	320	-6.003 -38.996 147.162 1.00 2		C	0
40	ATOM	5715	N	SER	321	-5.350 -40.177 145.366 1.00 2		С	N
40	ATOM	5716	CA	SER	321	-6.194 -41.336 145.675 1.00 2		С	С
	ATOM	5717	CB	SER	321	-6.100 -42.394 144.571 1.00 2		C	С
	MOTA	5718	OG	SER	321	-6.657 -41.916 143.365 1.00 2		C	
	ATOM	5719	C	SER	321	-5.757 -41.973 146.985 1.00 2		C	C
45	MOTA	5720	0	SER	321	-6.585 -42.420 147.766 1.00 2		C	0
45	ATOM	5721	N	ILE	322	-4.448 -42.031 147.208 1.00 2		С	N
	MOTA	5722	CA	ILE	322	-3.905 -42.605 148.428 1.00 2		C	C
	ATOM	5723	CB	ILE	322	-2.357 -42.707 148.336 1.00 2		C	C
	MOTA	5724		ILE	322	-1.741 -42.841 149.730 1.00 2		C	C
<b>F</b> 0	MOTA	5725		ILE	322	-1.980 -43.902 147.444 1.00 2		C	C
50	MOTA	5726		ILE	322	-0.504 -43.985 147.088 1.00 2		C	C
	MOTA	5727	C	ILE	322	-4.327 -41.754 149.629 1.00 2		С	C
	MOTA	5728	0	ILE	322	-4.642 -42.281 150.693 1.00 1		С	0
	MOTA	5729	N	ASN	323	-4.328 -40.437 149.442 1.00 2		C	N
EE	ATOM	5730	CA	ASN	323	-4.735 -39.483 150.478 1.00 2		C	С
55	ATOM	5731	CB	ASN	323	-4.665 -38.049 149.925 1.00		С	С
	ATOM	5732	CG	ASN	323	-5.148 -36.998 150.922 1.00		С	C
	MOTA	5733		LASN	323	-6.252 -37.087 151.463 1.00		С	0
	MOTA	5734	ND2	2 ASN	323	<b>-4.323 -35.986 151.151 1.00</b>	29.48	С	N

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	MOTA	5735	С	ASN	323		-39.813		1.00 22.69	С	С
	MOTA	5736	0	ASN	323		-39.936		1.00 22.78	C	0
	MOTA	5737	N	GLU	324	-7.025	-39.959	149.859	1.00 22.11	C	N
_	MOTA	5738	CA	GLU	324		-40.294		1.00 23.38	С	C
5	MOTA	5739	CB	GLU	324		-40.263		1.00 24.94	С	С
	ATOM	5740	CG	GLU	324		-38.866		1.00 28.85	C	С
	ATOM	5741	CD	GLU	324	-10.021			1.00 31.28	C	C
	MOTA	5742		GLU	324			146.498	1.00 34.04	С	0
	MOTA	5743	OE2	GLU	324			145.976	1.00 32.20	С	0
10	MOTA	5744	С	GLU	324			150.714	1.00 22.12	C	С
	MOTA	5745	0	GLU	324			151.575	1.00 21.79	С	0
	MOTA	5746	N	ALA	325	-7.781	-42.641	150.297	1.00 20.25	C	N
	MOTA	5747	CA	ALA	325			150.869	1.00 20.09	C	С
	MOTA	5748	CB	ALA	325			150.112	1.00 20.27	C	С
15	MOTA	5749	С	ALA	325			152.354	1.00 19.28	C	С
	MOTA	5750	0	ALA	325			153.155	1.00 20.38	С	0
	MOTA	5751	N	TYR	326			152.728	1.00 18.75	С	N
	MOTA	5752	CA	TYR	326			154.140	1.00 19.01	C	С
	MOTA	5753	CB	TYR	326	-5.147	-41.869	154.325	1.00 18.00	C	С
20	MOTA	5754	CG	TYR	326	-3.719	-42.355	154.417	1.00 18.20	С	C
	MOTA	5755	CD1	TYR	326			155.513	1.00 17.45	С	С
	MOTA	5756	CE1		326			155.625	1.00 15.87	С	С
	MOTA	5757	CD2	TYR	326	-2.793	-42.022	153.433	1.00 17.03	C	С
	ATOM	5758	CE2	TYR	326	-1.466	-42.411	153.536	1.00 16.49	C	C
25	MOTA	5759	CZ	TYR	326	-1.045	-43.148	154.633	1.00 16.11	С	C
	MOTA	5760	OH	TYR	326	0.281	-43.497	154.719	1.00 14.54	C	0
	MOTA	5761	С	TYR	326	-7.437	-42.601	154.936	1.00 19.02	С	C
	MOTA	5762	0	TYR	326	-7.673	-43.127	156.029	1.00 16.95	C	0
	MOTA	5763	N	GLY	327	-8.223	-41.676	154.382	1.00 18.76	С	N
30	MOTA	5764	CA	GLY	327	-9.448	-41.253	155.036	1.00 20.32	C	С
	MOTA	5765	С	GLY	327	-10.379	-42.434	155.263	1.00 21.04	С	С
	MOTA	5766	0	GLY	327	-10.973	-42.576	156.329	1.00 20.76	C	0
	ATOM	5767	N	TYR	328	-10.516	-43.289	154.257	1.00 21.82	C	N
	MOTA	5768	CA	TYR	328	-11.366	-44.463	154.398	1.00 22.78	С	C
35	MOTA	5769	CB	TYR	328	-11.461	-45.200	153.055	1.00 24.82	С	C
	ATOM	5770	CG	TYR	328			153.120	1.00 27.89	C	C
	MOTA	5771	CD1	TYR	328	-11.645	-47.675	153.544	1.00 28.22	С	C
	MOTA	5772	CE1	TYR	328			153.680	1.00 29.71	С	С
	ATOM	5773	CD2		328			152.823	1.00 29.15	С	C
40	MOTA	5774	CE2		328			152.959	1.00 30.37	C	C
	MOTA	5775	CZ	TYR	328			153.391	1.00 29.77	С	С
	MOTA	5776		TYR	328			153.567	1.00 31.94	С	
	MOTA	5777	С	TYR	328			155.495	1.00 22.29	С	С
	MOTA	5778	0	TYR	328			3 156.391	1.00 21.48	C	0
45	MOTA	5779	N	GLN	329			3 155.427	1.00 22.12	С	N
	MOTA	5780	CA	GLN	329			156.424	1.00 22.84	С	C
	MOTA	5781	CB	GLN	329			156.205	1.00 21.43	С	С
	MOTA	5782	CG	GLN	329			2 154.901	1.00 20.52	С	C
	MOTA	5783	CD	GLN	329			L 154.789	1.00 21.36	C	С
50	ATOM	5784		. GLN				3 153.889	1.00 21.96	C	0
	ATOM	5785		GLN	329			3 155.712	1.00 17.88	С	N
	MOTA	5786	С	GLN	329			3 157.846	1.00 24.04	C	С
	MOTA	5787	0	GLN	329			1 158.719	1.00 24.21	С	0
	MOTA	5788	N	ILE	330			3 158.070	1.00 24.50	С	N
55	MOTA	5789		ILE	330			9 159.382	1.00 26.20	С	С
	MOTA	5790	СВ	ILE	330			1 159.395		С	C
	MOTA	5791		2 ILE	330	-9.413	41.804	4 160.470	1.00 26.57	С	С
	MOTA	5792	CG1	LILE	330	-7.196	-42.562	2 159.670	1.00 28.06	С	С

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	MOTA	5793	CD1		330		-43.192		1.00 29.01	C	С
	MOTA	5794	С	ILE	330	-10.634			1.00 26.85	С	C
	ATOM	5795	0	ILE	330	-10.879			1.00 24.12	С	0
_	ATOM	5796	N	GLN	331	-11.594			1.00 28.57	C	N
5	MOTA	5797	CA	GLN	331	-12.987			1.00 31.36	C	С
	MOTA	5798	CB	GLN	331	-13.887	-43.474	158.389	1.00 34.12	С	С
	MOTA	5799	CG	GLN	331	-14.097			1.00 40.12	С	C
	MOTA	5800	CD	GLN	331	-15.111	-45.281	157.055	1.00 43.64	С	С
	MOTA	5801	OE1	GLN	331	-14.886	-46.346	156.463	1.00 46.08	C	0
10	MOTA	5802	NE2	GLN	331			157.737	1.00 45.76	C	N
	MOTA	5803	С	GLN	331			159.493	1.00 30.71	C	С
	MOTA	5804	0	GLN	331	-14.393	-45.995	160.220	1.00 30.97	С	0
	MOTA	5805	N	HIS	332	-12.778	-46.608	158.801	1.00 30.40	С	N
	MOTA	5806	CA	HIS	332	-13.154	-48.017	158.802	1.00 29.57	C	C
15	MOTA	5807	CB	HIS	332	-12.897	-48.605	157.411	1.00 30.43	С	С
	MOTA	5808	CG	HIS	332	-13.480	-49.969	157.209	1.00 32.56	C	С
	MOTA	5809	CD2	HIS	332	-12.898	-51.148	156.881	1.00 33.11	С	С
	MOTA	5810	ND1	HIS	332	-14.830	-50.225	157.314	1.00 33.92	С	N
	MOTA	5811	CE1	HIS	332	-15.055	-51.502	157.057	1.00 33.97	С	C
20	MOTA	5812	NE2	HIS	332	-13.899	-52.084	156.791	1.00 33.45	C	N
	MOTA	5813	C	HIS	332	-12.471	-48.881	159.861	1.00 28.67	С	С
	ATOM	5814	0	HIS	332	-13.054	-49.855	160.333	1.00 28.01	C	0
	ATOM	5815	N	ILE	333	-11.248	-48.532	160.247	1.00 27.55	С	N
	ATOM	5816	CA	ILE	333	-10.541	-49.336	161.237	1.00 26.64	С	С
25	ATOM	5817	CB	ILE	333	-9.264	-49.938	160.623	1.00 26.21	С	С
	ATOM	5818	CG2		333	-8.508	-50.761	161.662	1.00 26.49	С	С
	ATOM	5819	CG1	ILE	333	-9.653	-50.844	159.447	1.00 25.58	С	С
	MOTA	5820		ILE	333			158.209	1.00 24.86	С	С
	ATOM	5821	С	ILE	333	-10.211	-48.587	162.519	1.00 26.30	С	C
30	MOTA	5822	0	ILE	333			162.525	1.00 26.27	C	0
	ATOM	5823	N	GLN	334	-10.853	-49.009	163.605	1.00 25.96	C	N
	MOTA	5824	CA	GLN	334	-10.678	-48.412	164.929	1.00 26.54	С	С
	ATOM	5825	СВ	GLN	334	-11.595	-49.132	165.923	1.00 28.84	C	C
	ATOM	5826	CG	GLN	334			167.336	1.00 32.45	Ċ	C
35	ATOM	5827	CD	GLN	334	-12.537	-49.386	168.254	1.00 36.09	C	С
	ATOM	5828	OE1		334			169.198	1.00 37.52	C	0
	ATOM	5829	NE2		334			167.969	1.00 37.02	C	N
	ATOM	5830	С	GLN	334			165.394	1.00 25.97	C	C
	ATOM	5831	O	GLN	334		-49.547		1.00 25.42	Ċ	O
40	ATOM	5832	N	GLY	335			165.919	1.00 25.33	C	N
	ATOM	5833	CA	GLY	335			166.392	1.00 24.56	C	C
	MOTA	5834	C	GLY	335			165.465	1.00 24.72		C
	ATOM	5835	0	GLY	335			165.917	1.00 24.71	C	0
	ATOM	5836	N	LEU	336			164.166	1.00 24.12	С	N
45	ATOM	5837	CA	LEU	336			163.194	1.00 24.24	C	C
	MOTA	5838	СВ	LEU	336			3 161.771	1.00 23.67	Č	Ċ
	ATOM	5839	CG	LEU	336			3 161.229	1.00 25.15	Č	Ċ
	ATOM	5840		LEU	336			5 159.795	1.00 24.62	Č	C
	ATOM	5841		LEU	336			7 161.277	1.00 24.69	č	Ċ
50	ATOM	5842	C	LEU	336			5 163.390	1.00 23.69	c	Č
00	ATOM	5843	ŏ	LEU	336			2 163.348	1.00 23.11	c	ō
	ATOM	5844	N	SER	337			163.598	1.00 23.42	Č	N
	ATOM	5845	CA	SER	337			l 163.771	1.00 23.42	c	C
	ATOM	5846	CB	SER	337			l 164.044	1.00 23.77	C	C
55	ATOM	5847	OG	SER	337			5 165.303	1.00 27.27		0
	ATOM	5848	C	SER	337			164.889			C
	ATOM	5849	0	SER	337			7 164.824			0
	ATOM	5850		ALA	338			7 104.824 2 165.907			N
	ATOM	2020	TA	YUN.	220	-3.31	-42.00	. TOJ.30/	1.00 21.30	_	1.4

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	MOTA	5851	CA	ALA	338	-4.649	-42.568	167.038	1.00 21.23	С	С
	MOTA	5852	CB	ALA	338		-43.660		1.00 19.99	С	C
	ATOM	5853	С	ALA	338		-42.372		1.00 20.93	С	С
-	MOTA	5854	0	ALA	338		-41.788		1.00 20.73	C	0
5	MOTA	5855	N	MET	339		-42.862		1.00 21.18	C	N
	MOTA	5856	CA	MET	339		-42.709		1.00 21.93	С	С
	MOTA	5857	CB	MET	339		-43.866		1.00 20.73	С	С
	MOTA	5858	CG	MET	339		-45.183		1.00 19.60	C	С
40	MOTA	5859	SD	MET	339		-46.523		1.00 19.45	С	S
10	MOTA	5860	CE	MET	339		-46.789		1.00 19.70	С	С
	MOTA	5861	С	MET	339			164.335	1.00 22.95	C	С
	MOTA	5862	0	MET	339			163.945	1.00 21.85	С	0
	MOTA	5863	N	MET	340			164.167	1.00 24.72	C	N
4 =	ATOM	5864	CA	MET	340			163.568	1.00 27.94	C	С
15	ATOM	5865	CB	MET	340			162.207	1.00 27.04	С	C
	ATOM	5866	CG	MET	340			161.453	1.00 26.07	С	С
	MOTA	5867	SD	MET	340			161.038	1.00 25.64	C	S
	ATOM	5868	CE	MET	340			159.656	1.00 24.53	С	С
20	ATOM	5869	C	MET	340			164.558	1.00 31.10	C	С
20	ATOM	5870	0	MET	340			164.193	1.00 29.95	C	0
	ATOM	5871 5872	N	PRO	341			165.825	1.00 34.82	С	N
	ATOM		CD	PRO	341			166.110	1.00 34.96	C	C
	MOTA	5873 5874	CA CB	PRO	341			166.934 167.904	1.00 38.26	C	C
25	ATOM ATOM	5875	CG	PRO PRO	341 341				1.00 37.12	C	C
25	ATOM	5876	C	PRO	341			167.016 166.490	1.00 35.89	С	С
	MOTA	5877	0	PRO	341			165.531	1.00 41.84 1.00 43.72	C	C 0
	ATOM	5878	N	LEU	342			167.196	1.00 43.72	C	
	ATOM	5879	CA	LEU	342			166.723	1.00 44.96	C	N C
30	ATOM	5880	CB	LEU	342			167.708	1.00 48.46	C	C
00	ATOM	5881	CG	LEU	342			168.952	1.00 49.28	C	C
	ATOM	5882		LEU	342			169.995	1.00 49.28	C	C
	ATOM	5883		LEU	342			169.532	1.00 50.31	C	C
	ATOM	5884	C	LEU	342			165.437	1.00 47.75	Č	C
35	ATOM	5885	ō	LEU	342			164.857	1.00 46.19	c	ō
	ATOM	5886	N	LEU	343			164.982	1.00 48.81	c	N
	ATOM	5887	CA	LEU	343			163.767	1.00 49.66	C	C
	ATOM	5888	СВ	LEU	343			163.875	1.00 47.94	C	Č
	ATOM	5889	CG	LEU	343			164.841	1.00 46.64	C	C
40	MOTA	5890	CD1	LEU	343	0.840	-31.632	164.275	1.00 46.63	С	С
	MOTA	5891	CD2	LEU	343	-0.438	-32.523	166.230	1.00 46.74	С	С
	ATOM	5892	С	LEU	343	-3.196	-33.071	162.607	1.00 51.07	С	С
	ATOM	5893	0	LEU	343	-3.014	-34.222	162.192	1.00 50.79	С	0
	MOTA	5894	N	GLN	344			162.125	1.00 53.04	С	N
45	MOTA	5895	CA	GLN	344	-4.988	-32.595	160.988	1.00 55.20	С	С
	MOTA	5896	CB	GLN	344			159.782	1.00 55.64	С	С
	MOTA	5897	CG	GLN	344			159.666	1.00 56.26	С	C
	MOTA	5898	CD	GLN	344			158.953	1.00 57.17	С	C
	MOTA	5899		GLN	344			158.912	1.00 57.87	C	0
50	MOTA	5900		GLN	344			158.384	1.00 57.99	C	N
	ATOM	5901	С	GLN	344			161.207	1.00 56.43	С	C
	MOTA	5902	0	GLN	344			160.349	1.00 56.91	С	0
	ATOM	5903	N	GLU	345			162.343	1.00 57.98	С	Ŋ
EE	MOTA	5904	CA	GLU	345			162.680	1.00 58.92	C	C
55	MOTA	5905	CB	GLU	345			162.696	1.00 59.25	C	C
	MOTA	5906	CG	GLU	345			163.412	1.00 60.42	С	С
	MOTA	5907	CD	GLU	345			162.449	1.00 60.91	C	C
	MOTA	5908	OE1	GLU	345	-8.396	-31.491	162.807	1.00 61.74	С	0

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	ATOM	5909	OE2	CLII	345	-0.205	-32.905	161 226	1.00 61.02	~	0
	ATOM	5910	C	GLU	345			161.719	1.00 51.02	C	c
	ATOM	5911		GLU	345		-36.638		1.00 59.19		Ö
			0							C	
5	MOTA	5912	OXT	GLU	345	-7.281	-37.761	161.931	1.00 59.81	C	0
5	TER	5913		GLU	345					С	_
	ATOM	5914	CB	PRO	103	-18.301			1.00 85.60	D	C
	MOTA	5915	CG	PRO	103	-19.706			1.00 85.87	D	C
	MOTA	5916	C	PRO	103	-16.716			1.00 84.96	D	C
40	MOTA	5917	0	PRO	103	-16.171			1.00 85.09	D	0
10	MOTA	5918	N	PRO	103			124.910	1.00 85.59	D	N
	MOTA	5919	CD	PRO	103			124.343	1.00 85.67	D	С
	MOTA	5920	CA	PRO	103			125.986	1.00 85.36	D	С
	MOTA	5921	N .	VAL	104			124.722	1.00 84.12	D	N
	MOTA	5922	CA	VAL	104			124.383	1.00 83.16	D	C
15	MOTA	5923	CB	VAL	104			124.512	1.00 83.15	D	С
	MOTA	5924		VAL	104			124.170	1.00 83.06	D	С
	MOTA	5925		VAL	104			123.588	1.00 83.15	D	C
	MOTA	5926	С	VAL	104			125.235	1.00 82.38	D	С
	MOTA	5927	0	VAL	104	-13.000	-92.790	124.709	1.00 82.56	D	0
20	ATOM	5928	N	GLN	105			126.541	1.00 81.19	D	N
	ATOM	5929	CA	GLN	105	-12.769	-92.406	127.423	1.00 79.59	D	C
	MOTA	5930	CB	GLN	105			127.082	1.00 80.41	D	С
	MOTA	5931	CG	GLN	105			127.413	1.00 81.01	D	C
	MOTA	5932	CD	GLN	105	-10.121	-93.381	128.905	1.00 81.38	D	C
25	MOTA	5933	OE1	GLN	105	-9.699	-92.494	129.648	1.00 81.47	D	0
	MOTA	5934	NE2	GLN	105	-10.452	-94.591	129.348	1.00 81.27	D	N
	MOTA	5935	С	GLN	105	-13.029	-92.153	128.912	1.00 77.80	D	C
	MOTA	5936	0	GLN	105	-12.190	-91.579	129.607	1.00 77.77	D	0
	MOTA	5937	N	LEU	106	-14.196	-92.573	129.394	1.00 75.34	D	N
30	ATOM	5938	CA	LEU	106			130.806	1.00 72.86	D	C
	MOTA	5939	CB	LEU	106	-15.641	~91.358	131.007	1.00 72.85	D	C
	ATOM	5940	CG	LEU	106	-17.068	-91.595	130.507	1.00 72.73	D	C
	ATOM	5941		LEU	106	-17.971	-90.477	131.006	1.00 72.42	D	C
	ATOM	5942	CD2	LEU	106			128.995	1.00 72.91	D	C
35	MOTA	5943	C	LEU	106	-15.024	-93.774	131.322	1.00 70.93	D	С
	ATOM	5944	0	LEU	106	-16.207	-93.981	131.588	1.00 70.45	D	0
	MOTA	5945	N	SER	107	-14.074	-94.693	131.450	1.00 68.94	D	N
	MOTA	5946	CA	SER	107	-14.343	-96.045	131.910	1.00 67.00	D	C
	MOTA	5947	CB	SER	107	-13.023	-96.776	132.172	1.00 66.72	D	C
40	MOTA	5948	OG	SER	107	-12.247	-96.102	133.146	1.00 65.92	D	0
	MOTA	5949	С	SER	107			133.160	1.00 65.96	D	C
	ATOM	5950	0	SER	107	-15.463	-95.079	133.806	1.00 65.94	D	0
	MOTA	5951	N	LYS	108			133.489	1.00 64.35	D	N
	MOTA	5952	CA	LYS	108	-16.489	-97.493	134.669	1.00 62.78	D	C
45	MOTA	5953	CB	LYS	108	-17.282	-98.798	134.547	1.00 63.38	D	C
	MOTA	5954	CG	LYS	108	-18.220	-98.836	133.341	1.00 64.14	D	С
	MOTA	5955	CD	LYS	108			133.350	1.00 65.05	D	С
	MOTA	5956	CE	LYS	108			3 132.112	1.00 65.31	D	С
	MOTA	5957	NZ	LYS	108			2 130.858	1.00 65.55	D	N
50	ATOM	5958	С	LYS	108			135.917	1.00 61.20	D	C
	ATOM	5959	0	LYS	108	-16.086	-97.290	137.025	1.00 60.95	D	0
	ATOM	5960	N	GLU	109	-14.317	-97.787	7 135.731	1.00 59.39	D	N
	MOTA	5961	CA	GLU	109			2 136.840	1.00 57.92	D	С
	ATOM	5962	CB	GLU	109			136.392	1.00 58.83	D	C
55	MOTA	5963	CG	GLU	109			7 135.870	1.00 60.46	D	C
	MOTA	5964	CD	GLU	109			34.372	1.00 61.30	D	С
	MOTA	5965		. GLU	109			133.936	1.00 61.45	D	0
	MOTA	5966	OE2	GLU	109	-11.344	-100.228	3 133.626	1.00 61.74	D	0

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	MOTA	5967	С	GLU	109	-13.182	-96.405	137.372	1.00 56.28	D	С
	MOTA	5968	0	GLU	109	-13.311			1.00 56.26	D	0
	MOTA	5969	N	GLN	110	-12.872			1.00 53.97	D	N
_	MOTA	5970	CA	GLN	110	-12.669	-94.090	136.868	1.00 51.62	D	С
5	MOTA	5971	СВ	GLN	110	-11.900	-93.336	135.775	1.00 51.23	D	С
	ATOM	5972	CG	GLN	110	-12.575	-93.289	134.422	1.00 50.22	D	С
	MOTA	5973	CD	GLN	110	-11.624	-92.904	133.295	1.00 49.90	D	C
	MOTA	5974	OE1	GLN	110	-12.062			1.00 49.66	D	0
<u></u>	ATOM	5975	NE2	GLN	110	-10.320	-93.051	133.528	1.00 49.10	D	N
10	MOTA	5976	С	GLN	110	-14.002	-93.410	137.190	1.00 50.17	D	С
	MOTA	5977	0	GLN	110	-14.042	-92.403	137.893	1.00 49.22	D	0
	ATOM	5978	N	GLU	111	-15.098	-93.965	136.683	1.00 48.81	D	N
	MOTA	5979	CA	GLU	111	-16.415	-93.418	136.984	1.00 47.31	D	С
	MOTA	5980	CB	GLU	111		-93.987		1.00 49.14	D	С
15	MOTA	5981	CG	GLU	111		-93.276		1.00 51.36	D	C
	ATOM	5982	CD	GLU	111	-19.723	-93.664	134.914	1.00 53.11	D	С
	MOTA	5983	OE1	GLU	111	-20.068	-94.862	134.793	1.00 53.51	D	0
	MOTA	5984	OE2	GLU	111	-20.098	-92.765	134.126	1.00 53.59	D	0
	MOTA	5985	С	GLU	111			138.440	1.00 45.08	D	C
20	ATOM	5986	0	GLU	111			139.127	1.00 44.43	D	0
	ATOM	5987	N	GLU	112	-16.191	-94.919	138.896	1.00 42.54	D	N
	ATOM	5988	CA	GLU	112	-16.378	-95.401	140.263	1.00 40.31	D	С
	MOTA	5989	СВ	GLU	112	-16.100	-96.912	140.331	1.00 40.53	D	C
	MOTA	5990	CG	GLU	112	-15.884	-97.484	141.744	1.00 41.61	D	С
25	ATOM	5991	CD	GLU	112			142.629	1.00 41.94	D	C
	MOTA	5992	OE1	GLU	112	-17.006	-97.761	143.831	1.00 42.13	D	0
	MOTA	5993	OE2	GLU	112	-18.219	-97.081	142.129	1.00 41.78	D	0
	MOTA	5994	С	GLU	112			141.199	1.00 38.58	D	С
	MOTA	5995	0	GLU	112			142.358	1.00 37.73	D	0
30	MOTA	5996	N	LEU	113			140.690	1.00 36.41	D	N
	MOTA	5997	CA	LEU	113			141.469	1.00 34.41	D	С
	MOTA	5998	CB	LEU	113			140.660	1.00 34.39	D	C
	MOTA	5999	CG	LEU	113			141.214	1.00 34.64	D	С
	ATOM	6000		LEU	113			142.556	1.00 33.12	D	С
35	MOTA	6001		LEU	113			140.200	1.00 34.83	D	C
	MOTA	6002	C	LEU	113			141.840	1.00 33.24	D	C
	MOTA	6003	0	LEU	113			142.961	1.00 32.72	D	0
	MOTA	6004	N	ILE	114			140.881	1.00 31.70	D	N
4.0	MOTA	6005	CA	ILE	114			141.109	1.00 31.14	D	C
40	MOTA	6006	CB	ILE	114			139.794	1.00 30.51	D	C
	MOTA	6007	CG2	ILE	114			140.087	1.00 28.60	D	С
	ATOM	6008		. ILE	114			138.870	1.00 30.78	D	С
	ATOM	6009		LE	114			137.532	1.00 29.75	D	С
45	MOTA	6010	C	ILE	114			142.140	1.00 31.12	D	C
45	MOTA	6011	0	ILE	114			143.015	1.00 30.80	D	0
	MOTA	6012	N	ARG	115			142.035	1.00 30.77	D	N
	MOTA	6013	CA	ARG	115			142.968	1.00 30.08	D	С
	ATOM	6014	CB	ARG	115			142.640	1.00 31.12	D	C
50	MOTA	6015	CG	ARG	115			143.393	1.00 32.38	D	C
50	MOTA	6016	CD	ARG	115			143.216	1.00 33.88	D	С
	ATOM	6017	NE	ARG	115			143.907	1.00 34.78	D	N
	ATOM	6018	CZ	ARG	115			145.232	1.00 35.85	D	С
	ATOM	6019		LARG	115			146.048		D	N
EE	ATOM	6020		2 ARG	115			145.746		D	N
55	ATOM	6021	C	ARG	115			144.393		D	С
	MOTA	6022		ARG	115			2 145.301		D	0
	MOTA	6023		THR	116			144.559		D	N
	MOTA	6024	CA	THR	116	-15.850	92.70	5 145.837	1.00 27.53	D	C

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	MOTA	6025		THR	116	-14.748	-93.769	145.676	1.00 28.08	D	С
	MOTA	6026		THR	116	-15.353			1.00 30.70	D	0
	MOTA	6027	CG2	THR	116	-13.978	-93.964	146.978	1.00 29.30	D	С
_	MOTA	6028	С	THR	116	-15.233			1.00 26.60	D	C
5	MOTA	6029	0	THR	116	-15.435			1.00 25.28	D	0
	MOTA	6030	N	LEU	117	-14.467	-90.715	145.622	1.00 24.99	D	N
	MOTA	6031	CA	LEU	117	-13.831			1.00 23.90	D	C
	MOTA	6032	CB	LEU	117	-12.951	-88.902	144.969	1.00 22.02	D	С
	MOTA	6033	CG	LEU	117	-11.621	-89.624	144.740	1.00 20.36	D	С
10	ATOM	6034	CD1	LEU	117	-10.988	-89.149	143.448	1.00 19.86	D	С
	MOTA	6035	CD2	LEU	117	-10.698	-89.379	145.920	1.00 19.89	D	С
	MOTA	6036	С	LEU	117	-14.882	-88.469	146.493	1.00 23.81	D	С
	MOTA	6037	0	LEU	117	-14.759	-87.823	147.533	1.00 23.44	D	0
	MOTA	6038	N	LEU	118	-15.920	-88.345	145.668	1.00 23.97	D	N
15	MOTA	6039	CA	LEU	118	-17.008	-87.409	145.908	1.00 24.68	D	С
	ATOM	6040	CB	LEU	118	-17.918	-87.350	144.680	1.00 26.07	D	С
	MOTA	6041	CG	LEU	118	-18.967	-86.236	144.665	1.00 27.99	D	С
	MOTA	6042	CD1	LEU	118	-18.298	-84.877	144.878	1.00 28.05	D	С
	MOTA	6043	CD2	LEU	118	-19.703	-86.266	143.329	1.00 29.41	D	С
20	MOTA	6044	С	LEU	118	-17.825	-87.775	147.144	1.00 24.04	D	С
	MOTA	6045	0	LEU	118	-18.187	-86.907	147.936	1.00 23.27	D	0
	MOTA	6046	N	GLY	119	-18.128	-89.061	147.295	1.00 23.56	D	N
	ATOM	6047	CA	GLY	119	-18.886	-89.501	148.451	1.00 22.61	D	С
	ATOM	6048	C	GLY	119	-18.142	-89.112	149.714	1.00 22.12	D	С
25	MOTA	6049	0	GLY	119	-18.715	-88.489	150.610	1.00 21.52	D	0
	MOTA	6050	N	ALA	120	-16.860	-89.471	149.776	1.00 21.08	D	N
	MOTA	6051	CA	ALA	120	-16.025	-89.163	150.931	1.00 21.42	D	C
	ATOM	6052	CB	ALA	120	-14.643	-89.795	150.761	1.00 20.72	D	С
	ATOM	6053	С	ALA	120	-15.886	-87.657	151.131	1.00 21.67	D	C
30	ATOM	6054	0	ALA	120	-15.933	-87.158	152.251	1.00 21.20	D	0
	ATOM	6055	N	HIS	121			150.037	1.00 21.91	D	N
	ATOM	6056	CA	HIS	121	-15.563	-85.493	150.118	1.00 22.79	D	С
	ATOM	6057	CB	HIS	121	-15.227	-84.923	148.734	1.00 23.76	D	С
	MOTA	6058	CG	HIS	121			148.680	1.00 24.65	D	С
35	ATOM	6059	CD2	HIS	121	-14.268	-82.525	148.980	1.00 25.00	D	С
	ATOM	6060	ND1	HIS	121	-16.337	-82.700	148.300	1.00 25.79	D	N
	ATOM	6061	CE1	HIS	121	-16.057	-81.410	148.366	1.00 25.43	D	C
	ATOM	6062	NE2	HIS	121	-14.807	-81.277	148.777	1.00 26.28	D	N
	ATOM	6063	С	HIS	121	-16.823	-84.834	150.674	1.00 22.33	D	C
40	ATOM	6064	0	HIS	121			151.557	1.00 21.50	D	0
	ATOM	6065	N	THR	122	-17.977	-85.234	150.156	1.00 21.62	D	N
	ATOM	6066	CA	THR	122	-19.249	-84.673	150.597	1.00 22.81	D	С
	ATOM	6067	CB	THR	122			149.772	1.00 23.18	D	C
	ATOM	6068	OG1		122			148.387	1.00 25.50	D	0
45	ATOM	6069	CG2		122			150.212	1.00 22.28	D	C
	ATOM	6070	С	THR	122			152.077	1.00 22.45	D	C
	ATOM	6071	0	THR	122			152.796	1.00 21.90	D	0
	ATOM	6072	N	ARG	123			152.527	1.00 21.68	D	N
	MOTA	6073	CA	ARG	123			153.922	1.00 22.61	D	С
50	ATOM	6074	CB	ARG	123			154.149	1.00 22.11	D	Č
	ATOM	6075	CG	ARG	123			153.500	1.00 24.00	D	C
	ATOM	6076	CD	ARG	123			153.975	1.00 23.54	D	C
	ATOM	6077	NE	ARG	123			153.438	1.00 24.44	D	N
	ATOM	6078	CZ	ARG	123			152.207	1.00 25.95	D	C
55	ATOM	6079		ARG	123			151.356	1.00 24.95	D	N
_	ATOM	6080		ARG	123			151.828		D	N
	ATOM	6081		ARG	123			2 154.914		Ď	C
	ATOM	6082		ARG	123			155.929	1.00 22.70	D	0
			-							_	_

	ATOM	6083	N	HIS	124	-17.318	-85.634	154.604	1.00 22.37	D	N
	ATOM	6084	CA	HIS	124	-16.359			1.00 22.99	D	C
	ATOM	6085	СВ	HIS	124	-15.223			1.00 23.32	D	Ċ
	MOTA	6086	CG	HIS	124	-15.699			1.00 25.33	D	C
5	ATOM	6087	CD2		124	-16.458			1.00 24.81	D	Ċ
•	MOTA	6088		HIS	124	-15.444			1.00 25.61	D	N
	MOTA	6089		HIS	124	-16.028			1.00 25.40	D	C
	ATOM	6090		HIS	124	-16.650			1.00 24.90	D	N
	ATOM	6091	C	HIS	124		-83.643		1.00 22.18	D	c
10	ATOM	6092	0	HIS	124		-82.975		1.00 22.39	D	Ö
. •	ATOM	6093	N	MET	125		-83.202		1.00 21.09	Ď	N
	ATOM	6094	CA	MET	125		-81.902		1.00 21.77	D	c
	ATOM	6095	СВ	MET	125		-82.093		1.00 21.06	D	č
	MOTA	6096	CG	MET	125		-82.995		1.00 21.00	D	c
15	ATOM	6097	SD	MET	125		-83.007		1.00 24.77	D	s
	MOTA	6098	CE	MET	125			151.192	1.00 24.77	D	c
	MOTA	6099	C	MET	125			153.128	1.00 21.17	D	c
	MOTA	6100	ō	MET	125			153.549	1.00 21.24	D	ŏ
	ATOM	6101	N	GLY	126			152.104	1.00 21.62	D	N
20	ATOM	6102	CA	GLY	126			151.364	1.00 22.06	D	C
20	ATOM	6103	C	GLY	126			152.113	1.00 23.28	D	Č
	ATOM	6104	ŏ	GLY	126			151.619	1.00 23.14	D	ŏ
	ATOM	6105	N	THR	127			153.296	1.00 23.14	D	N
	ATOM	6106	CA	THR	127			154.071	1.00 24.53	D	c
25	ATOM	6107	CB	THR	127			154.170	1.00 25.29	D	C
20	ATOM	6108	OG1		127			154.456	1.00 25.66	D	õ
	ATOM	6109	CG2		127			152.856	1.00 25.86	D	č
	ATOM	6110	C	THR	127			155.473	1.00 24.35	D	c
	MOTA	6111	Ö	THR	127			156.387	1.00 24.95	D	ŏ
30	ATOM	6112	N	MET	128			155.651	1.00 23.31	D	N
00	ATOM	6113	CA	MET	128			156.969	1.00 23.31	D	C
	MOTA	6114	CB	MET	128			156.981	1.00 22.55	D	C
	ATOM	6115	CG	MET	128			156.440	1.00 19.76	D	č
	ATOM	6116	SD	MET	128			156.386	1.00 13.70	D	s
35	ATOM	6117	CE	MET	128			155.196	1.00 18.78	D	c
00	ATOM	6118	C	MET	128			157.437	1.00 21.87	D	Č
	ATOM	6119	Ö	MET	128			158.635	1.00 20.88	D	Ö
	ATOM	6120	N	PHE	129			156.496	1.00 20.44	D	N
	ATOM	6121	CA	PHE	129			156.822	1.00 22.40	D	c
40	ATOM	6122	СВ	PHE	129			155.521	1.00 23.38	D	Č
	ATOM	6123	CG	PHE	129			154.716	1.00 24.91	D	Č
	ATOM	6124	CD1		129			155.076	1.00 26.36	D	č
	ATOM	6125		PHE	129			153.608	1.00 26.00	D	Č
	ATOM	6126		PHE	129			154.344	1.00 27.45	D	Č
45	MOTA	6127		PHE	129			152.869	1.00 26.23	D	Č
. •	ATOM	6128	CZ	PHE	129			153.239	1.00 26.91	D	C
	MOTA	6129	C	PHE	129			157.682	1.00 22.06	D	č
	ATOM	6130	Ö	PHE	129			158.394	1.00 20.89	D	ō
	ATOM	6131	N	GLU	130			157.625	1.00 23.03	D	N
50	ATOM	6132	CA	GLU	130			158.416	1.00 24.67	D	C
00	MOTA	6133	СВ	GLU	130			157.978	1.00 26.82	D	C
	MOTA	6134	CG	GLU	130			156.573	1.00 29.47	D	C
	MOTA	6135	CD	GLU	130			156.467	1.00 32.46	D	c
	ATOM	6136		L GLU	130			7 157.480	1.00 34.27	D	0
55	ATOM	6137		GLU	130			155.362	1.00 33.81	D	o
	ATOM	6138	C	GLU	130			159.916	1.00 33.81	D	C
	ATOM	6139	ŏ	GLU	130			160.745	1.00 24.44	D	0
	ATOM	6140	N	GLN	131			3 160.743	1.00 22.76	D	N
				J		_3.507	, 3 , 50,	, 100.231	1.00 22.70	J	TA

	ATOM	6141	CA	GLN	131	-18.589	-75.853	161.652	1.00 22.20	D	С
	MOTA	6142	CB	GLN	131	-17.967	-77.245	161.778	1.00 24.32	D	С
	MOTA	6143	CG	GLN	131	-18.846	-78.385	161.290	1.00 28.51	D	С
_	ATOM	6144		GLN	131	-20.204			1.00 31.63	D	C
5	MOTA	6145	OE1	GLN	131	-20.305	-78.373	163.193	1.00 34.51	D	0
	ATOM	6146	NE2	GLN	131	-21.259	-78.420	161.154	1.00 32.89	D	N
	MOTA	6147	С	GLN	131	-17.627	-74.843	162.269	1.00 20.71	D	С
	MOTA	6148	0	GLN	131	-17.440	-74.829	163.483	1.00 19.33	D	0
	MOTA	6149	N	PHE	132		-74.007		1.00 18.82	D	N
10	MOTA	6150	CA	PHE	132	-16.055	-73.025	161.939	1.00 18.21	D	C
	MOTA	6151	CB	PHE	132	-15.547	-72.129	160.798	1.00 17.25	D	C
	ATOM	6152	CG	PHE	132	-14.612	-72.825	159.823	1.00 16.76	D	С
	MOTA	6153	CD1	PHE	132	-14.178	-74.133	160.039	1.00 15.85	D	С
	MOTA	6154	CD2	PHE	132	-14.151	-72.150	158.694	1.00 15.94	D	С
15	MOTA	6155	CE1	PHE	132	-13.299	-74.762	159.141	1.00 16.23	D	С
	ATOM	6156	CE2	PHE	132	-13.268	-72.772	157.789	1.00 16.64	D	С
	MOTA	6157	CZ	PHE	132	-12.843	~74.078	158.017	1.00 15.57	D	C
	ATOM	6158	С	PHE	132	-16.641	-72.159	163.050	1.00 17.13	D	С
	ATOM	6159	0	PHE	132	-15.946	-71.802	163.993	1.00 16.35	D	0
20	MOTA	6160	N	VAL	133	-17.921	-71.832	162.942	1.00 16.80	D	N
	MOTA	6161	CA	VAL	133	-18.584	-71.011	163.949	1.00 18.67	D	С
	MOTA	6162	СВ	VAL	133	-20.010	-70.597	163.480	1.00 18.65	D	С
	ATOM	6163	CG1	VAL	133	-20.978	-71.762	163.626	1.00 18.17	D	C
	ATOM	6164	CG2	VAL	133	-20.484	-69.400	164.272	1.00 21.43	D	С
25	ATOM	6165	С	VAL	133			165.327	1.00 18.01	D	C
	ATOM	6166	0	VAL	133	-19.034	-71.058	166.314	1.00 17.64	D	O
	ATOM	6167	N	GLN	134			165.383	1.00 17.24	D	N
	MOTA	6168	CA	GLN	134			166.631	1.00 18.01	D	C
	ATOM	6169	СВ	GLN	134			166.332	1.00 18.24	D	C
30	ATOM	6170	CG	GLN	134			165.643	1.00 20.23	D	C
	ATOM	6171	CD	GLN	134			166.579	1.00 20.77	D	C
	MOTA	6172	OE1		134			166.899	1.00 22.88	D	ō
	ATOM	6173	NE2		134			167.032	1.00 20.71	D	N
	ATOM	6174	С	GLN	134			167.390	1.00 18.05	D	C
35	ATOM	6175	Ō	GLN	134			168.514	1.00 17.44	D	ō
	ATOM	6176	N	PHE	135			166.778	1.00 17.41	D	N
	ATOM	6177	CA	PHE	135			167.392	1.00 17.91	D	C
	ATOM	6178	СВ	PHE				166.412	1.00 16.42	D	c
	ATOM	6179	CG	PHE	135			166.091	1.00 15.86	D	Ċ
40	ATOM	6180		PHE	135			166.982	1.00 16.41	D	Č
. •	ATOM	6181		PHE	135			164.942	1.00 15.95	D	Č
	ATOM	6182		PHE	135			166.737	1.00 17.52	D	C
	MOTA	6183	CE2	PHE	135	-14.756	-76.563	164.678	1.00 16.49	D	Č
	MOTA	6184	CZ	PHE	135			165.581	1.00 16.50	D	Č
45	ATOM	6185	C	PHE	135			167.909	1.00 18.76	Ď	C
	MOTA	6186	Ö	PHE	135			167.703	1.00 19.28	Ď	Ö
	ATOM	6187	N	ARG	136			168.590	1.00 19.27	D	Ŋ
	ATOM	6188	CA	ARG	136			169.198	1.00 20.81	Ď	C
	ATOM	6189	CB	ARG	136			170.428	1.00 22.56	D	Ċ
50	MOTA	6190	CG	ARG	136			171.538	1.00 26.27	D	C
00	ATOM	6191	CD	ARG	136			172.275	1.00 20.27	D	c
	MOTA	6192	NE	ARG	136			171.601	1.00 34.44	D	N
	ATOM	6193	CZ	ARG	136			171.612	1.00 34.44	D	C
	MOTA	6194		ARG	136			171.012	1.00 36.40	D	И
55	MOTA	6195		ARG	136			172.200	1.00 37.87	D	
	MOTA	6196	C	ARG				168.261	1.00 39.01	ם	N
	MOTA	6197	Ö	ARG				168.201	1.00 20.52	D	C
	ATOM	6198	N	PRO	137			168.529		D	0
	A LON	0190	7.4	LINO	13/	-13.207	-00.10/	TO1.T2T	1.00 20.21	ע	N

	MOTA	6199	CD	PRO	137	-16.587			1.00 20.40	D	С
	MOTA	6200	CA	PRO	137	-14.745			1.00 21.02	D	C
	MOTA	6201	СВ	PRO	137	-15.646		165.023	1.00 20.72	D	C
_	MOTA	6202	CG	PRO	137	-16.973			1.00 21.45	D	C
5	MOTA	6203	C	PRO	137	-14.806			1.00 21.01	D	С
	MOTA	6204	0	PRO	137	-15.789			1.00 20.39	D	0
	MOTA	6205	N	PRO	138	-13.735			1.00 21.11	D	N
	ATOM	6206	CD	PRO	138	-12.375		166.286	1.00 21.05	D	С
40	ATOM	6207	CA	PRO	138	-13.764			1.00 21.01	D	С
10	MOTA	6208	CB	PRO	138	-12.400			1.00 21.60	D	С
	MOTA	6209	CG	PRO	138	-11.523			1.00 22.95	D	С
	MOTA	6210	С	PRO	138		-62.830		1.00 20.24	D	С
	MOTA	6211	0	PRO	138		-63.174		1.00 17.44	D	0
4-	MOTA	6212	N	ALA	139		-61.804		1.00 19.19	D	N
15	MOTA	6213	CA	ALA	139		-60.991		1.00 17.76	D	С
	MOTA	6214	CB	ALA	139		-59.892		1.00 17.06	D	С
	MOTA	6215	С	ALA	139		-60.360		1.00 17.86	D	С
	ATOM	6216	0	ALA	139		-60.281		1.00 17.93	Ø	0
	MOTA	6217	N	HIS	140			165.130	1.00 17.55	D	N
20	MOTA	6218	CA	HIS	140			163.842	1.00 16.71	D	C
	ATOM	6219	CB	HIS	140			163.825	1.00 16.46	D	C
	MOTA	6220	CG	HIS	140			163.533	1.00 15.67	D	С
	MOTA	6221		HIS	140			162.371	1.00 14.99	D	С
	MOTA	6222	ND1	HIS	140	-11.745	-60.257	164.508	1.00 13.91	D	N
25	MOTA	6223		HIS	140			163.956	1.00 14.59	D	С
	MOTA	6224		HIS	140			162.661	1.00 15.09	D	N
	MOTA	6225	С	HIS	140	-15.013	-60.212	162.650	1.00 16.75	D	С
	MOTA	6226	0	HIS	140	-15.105	-59.764	161.513	1.00 15.99	D	0
	MOTA	6227	N	LEU	141.	-15.033	-61.518	162.901	1.00 17.79	D	N
30	MOTA	6228	CA	LEU	141			161.814	1.00 18.75	D	С
	ATOM	6229	CB	LEU	141	-14.682	-63.869	162.273	1.00 18.44	D	С
	ATOM	6230	CG	LEU	141			162.573	1.00 18.91	D	C
	MOTA	6231	CD1	LEU	141	-12.904	-65.504	162.883	1.00 17.51	D	С
	MOTA	6232	CD2	LEU	141	-12.345	-63.588	161.362	1.00 17.97	D	C
35	MOTA	6233	С	LEU	141			161.283	1.00 20.14	D	C
	MOTA	6234	0	LEU	141	-16.806	-63.055	160.167	1.00 19.48	D	0
	MOTA	6235	N	PHE	142			162.084	1.00 21.14	D	N
	MOTA	6236	CA	PHE	142			161.649	1.00 23.17	D	C
	MOTA	6237	CB	PHE	142			162.837	1.00 22.30	D	C
40	MOTA	6238	CG	PHE	142			163.726	1.00 22.33	D	С
	MOTA	6239		PHE	142			164.673	1.00 22.28	D	C
	MOTA	6240	CD2	PHE	142			163.578	1.00 22.17	D	C
	MOTA	6241		PHE	142			165.455	1.00 22.00	D	C
	MOTA	6242		PHE	142			164.353	1.00 20.89	D	C
45	MOTA	6243	CZ	PHE	142	-20.116	-65.550	165.293	1.00 21.02	D	C
	MOTA	6244	C	PHE	142			160.580	1.00 25.02	D	С
	MOTA	6245	0	PHE	142			160.692	1.00 24.19	D	0
	MOTA	6246	N	ILE	143			159.534	1.00 28.08	D	N
	MOTA	6247	CA	ILE	143	-20.281	-60.755	158.410	1.00 31.14	D	С
50	MOTA	6248	СВ	ILE	143			157.446	1.00 32.88	D	С
	MOTA	6249		ILE	143			157.814	1.00 33.01	D	С
	MOTA	6250		ILE	143			155.986	1.00 34.33	D	С
	MOTA	6251		ILE	143	-21.350	-59.741	155.606	1.00 36.85	D	С
	MOTA	6252	С	ILE	143			158.913	1.00 32.38	D	C
55	MOTA	6253	0	ILE	143			159.820	1.00 31.70	D	0
	MOTA	6254	N	HIS	144			158.334	1.00 33.55	D	N
	MOTA	6255		HIS	144			158.680		D	С
	MOTA	6256	CB	HIS	144	-22.543	-56.993	158.683	1.00 37.72	D	

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	MOTA	6257		HIS	144			С
	MOTA	6258	CD2		144			С
	MOTA	6259	ND1		144			N
_	MOTA	6260	CE1		144	·	D	С
5	MOTA	6261	NE2	HIS	144	<b>-24.186 -58.036 155.569 1.00 42.04</b>	D	N
	MOTA	6262	С	HIS	144	-20.491 -56.424 159.990 1.00 34.54	D	С
	MOTA	6263	0	HIS	144	-20.892 -55.329 160.386 1.00 34.20	D	0
	MOTA	6264	N	HIS	145	-19.605 -57.147 160.666 1.00 33.54	D	N
	ATOM	6265	CA	HIS	145	-19.048 -56.650 161.916 1.00 32.95	D	С
10	MOTA	6266	СВ	HIS	145			С
	ATOM	6267	CG	HIS	145			С
	ATOM	6268	CD2		145		D	Ċ
	ATOM	6269	ND1		145			N
	ATOM	6270	CE1		145		D	C
15	ATOM	6271	NE2	HIS	145		D	N
	ATOM	6272	C	HIS	145		D	C
	ATOM	6273	Ö	HIS	145		D	Ö
	ATOM	6274	N	GLN	146			
	ATOM	6275	CA		146		D	N
20				GLN			D	С
20	ATOM	6276	CB	GLN	146		D	C
	MOTA	6277	CG	GLN	146		D	C
	MOTA	6278	CD	GLN	146		D	С
	MOTA	6279		GLN	146		D	0
05	MOTA	6280	NE2	GLN	146	-16.870 -50.503 164.862 1.00 42.83	D	N
25	MOTA	6281	С	GLN	146	-15.087 -54.727 162.374 1.00 30.70	D	C
	MOTA	6282	0	GLN	146	-14.985 -55.716 163.093 1.00 29.52	D	0
	MOTA	6283	N	PRO	147	-14.067 -54.307 161.599 1.00 29.06	D	N
	MOTA	6284	CD	PRO	147	-14.144 -53.170 160.666 1.00 29.28	D	C
	MOTA	6285	CA	PRO	147	-12.738 -54.946 161.532 1.00 27.41	D	C
30	MOTA	6286	CB	PRO	147	-11.969 -54.050 160.559 1.00 27.81	D	С
	MOTA	6287	CG	PRO	147	-13.042 -53.488 159.682 1.00 28.67	D	С
	MOTA	6288	С	PRO	147	-12.040 -55.020 162.894 1.00 25.53	D	С
	MOTA	6289	0	PRO	147	-12.434 -54.328 163.831 1.00 26.05	D	0
	ATOM	6290	N	LEU	148	-11.008 -55.854 163.009 1.00 23.12	D	N
35	ATOM	6291	CA.	LEU	148	-10.282 -55.979 164.280 1.00 21.13	D	С
	ATOM	6292	СВ	LEU	148	-9.302 -57.153 164.231 1.00 20.59	D	C
	ATOM	6293	CG	LEU	148	-8.531 -57.403 165.532 1.00 20.63	D	Č
	ATOM	6294	CD1	LEU	148	-9.478 -57.969 166.595 1.00 18.94	D	Ċ
	ATOM	6295		LEU	148	-7.384 -58.372 165.270 1.00 20.92	D	Ċ
40	ATOM	6296	С	LEU	148	-9.509 -54.687 164.547 1.00 19.93	D	Ċ
	ATOM	6297	Ō	LEU	148	-8.641 -54.304 163.766 1.00 19.73	D	ŏ
	ATOM	6298	N	PRO	149	-9.815 -53.995 165.654 1.00 18.90	D	N
	ATOM	6299	CD	PRO	149	-10.844 -54.314 166.658 1.00 18.79	D	C
	ATOM	6300	CA	PRO	149	-9.129 -52.738 165.992 1.00 18.86	D	c
45	ATOM	6301	СВ	PRO	149	-9.726 -52.368 167.351 1.00 17.91	D	Ċ
10	MOTA	6302	CG	PRO	149	-11.102 -52.954 167.290 1.00 18.09	D	
	MOTA	6303	C	PRO	149	-7.608 -52.836 166.056 1.00 18.30		C
	MOTA	6304	Ö	PRO	149	-7.045 -53.894 166.310 1.00 18.17	D D	С
	ATOM	6305	N	THR	150	-6.954 -51.706 165.837 1.00 19.06		0
50							D	N
30	MOTA	6306	CA	THR	150	-5.509 -51.619 165.870 1.00 18.45	D	С
	ATOM	6307	CB	THR	150	-5.060 -50.141 165.785 1.00 18.97	D	С
	ATOM	6308		THR	150	-5.334 -49.633 164.469 1.00 16.80	D	0
	MOTA	6309	CG2		150	-3.574 -50.007 166.110 1.00 16.34	D	С
66	MOTA	6310	С	THR	150	-4.875 -52.247 167.112 1.00 19.56	D	С
55	ATOM	6311	0	THR	150	-3.923 -53.016 167.003 1.00 20.79	D	0
	MOTA	6312	N	LEU	151	-5.391 -51.928 168.292 1.00 19.20	D	N
	MOTA	6313	CA	LEU	151	-4.792 -52.451 169.518 1.00 19.49	D	С
	MOTA	6314	СВ	LEU	151	-4.714 -51.338 170.579 1.00 19.37	D	С

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	ATOM	6315	CG	LEU	151	-3.833	-50.116	170.260	1.00 19.71	D	С
	ATOM	6316	CD1		151	-3.960	-49.078	171.376	1.00 19.45	D	С
	ATOM	6317	CD2		151		-50.542		1.00 19.57	D	С
	ATOM	6318		LEU	151		-53.680		1.00 19.39	D	Č
5	ATOM	6319		LEU	151		-54.136		1.00 18.10	D	Ō
	ATOM	6320		ALA	152		-54.220		1.00 18.60	D	N
	ATOM	6321	CA	ALA	152		-55.396		1.00 18.53	D	C
	ATOM	6322	СВ	ALA	152		-55.695		1.00 17.95	D	Ċ
	ATOM	6323	C	ALA	152			169.980	1.00 18.77	D	Ċ
10	ATOM	6324	Ö	ALA	152			169.034	1.00 18.21	D	ō
. •	ATOM	6325	N	PRO	153			171.050	1.00 18.94	D	N
	ATOM	6326	CD	PRO	153			172.306	1.00 18.91	D	C
	ATOM	6327	CA	PRO	153			171.086	1.00 19.14	D	Ċ
	ATOM	6328	СВ	PRO	153			172.442	1.00 19.09	D	Ċ
15	ATOM	6329	CG	PRO	153			173.283	1.00 20.56	D	Ċ
.0	MOTA	6330	C	PRO	153			169.939	1.00 18.81	D	c
	ATOM	6331	Ö	PRO	153			169.589	1.00 17.99	D	ŏ
	ATOM	6332	N	VAL	154			169.350	1.00 18.43	D	N
	MOTA	6333	CA	VAL	154			168.269	1.00 18.36	D	c
20	ATOM	6334	CB	VAL	154			167.166	1.00 18.03	D	c
20	ATOM	6335		VAL	154			166.551	1.00 10.03	D	c
	ATOM	6336		VAL	154			167.747	1.00 17.94	D	Ċ
	MOTA	6337	C	VAL	154			168.787	1.00 17.34	D	c
	ATOM	6338	0	VAL	154			168.047	1.00 16.70	D	Ö
25	ATOM	6339	N	LEU	155			170.053	1.00 10.70	D	N
23	ATOM	6340	CA	LEU	155			170.637	1.00 17.75	D	C
	ATOM	6341	CB	LEU	155			172.156	1.00 18.87	D	C
	ATOM	6342	CG	LEU	155			172.130	1.00 18.81	D	c
	ATOM	6343		LEU	155			172.393	1.00 18.11	D	Ċ
30	ATOM	6344		LEU	155			174.426	1.00 19.81	D	c
30	ATOM	6345	CDZ	LEU	155			170.305	1.00 13.81	D	C
	ATOM	6346	0	LEU	155			169.897	1.00 10.45	D	Ö
	MOTA	6347	N	PRO	156			170.480	1.00 13.25	D	N
	ATOM	6348	CD	PRO	156			171.115	1.00 10.23	D	C
35	MOTA	6349	CA	PRO	156			170.153	1.00 17.20	D	c
00	ATOM	6350	CB	PRO	156			170.409	1.00 17.77	D	c
	MOTA	6351	CG	PRO	156			171.560	1.00 17.51	D	C
	MOTA	6352	C	PRO	156			168.714	1.00 17.31	D	C
	ATOM	6353	Ö	PRO	156			168.458	1.00 17.24	D	õ
40	ATOM	6354	N	LEU	157			167.780	1.00 16.35	D	N
40	ATOM	6355	CA	LEU	157			166.374	1.00 15.67	D	C
	ATOM	6356	CB	LEU	157			165.510	1.00 15.48	D	Č
	ATOM	6357	CG	LEU	157			164.007	1.00 14.81	D	C
	MOTA	6358		LEU	157			163.411	1.00 13.61	D	C
45	ATOM	6359		LEU	157			163.337	1.00 14.21	Ď	c
	ATOM	6360	C	LEU	157			166.151	1.00 15.16	D	Č
	ATOM	6361	Ö	LEU	157			165.405	1.00 14.60	D	Õ
	MOTA	6362	N	VAL	158			166.788	1.00 14.93	D	N
	ATOM	6363	CA	VAL	158			166.677	1.00 16.52	D	C
50	MOTA	6364	СВ	VAL	158			167.478	1.00 17.32	D	Č
00	MOTA	6365		VAL	158			167.538	1.00 16.08	D	c
	MOTA	6366		VAL	158			166.825	1.00 15.99	D	C
	ATOM	6367	C	VAL	158			2 167.213	1.00 16.29	D	
	ATOM	6368		VAL	158			166.604	1.00 14.70	D	0
55	ATOM	6369		THR	159			3 168.350	1.00 14.70	D	
50	ATOM	6370		THR	159			168.959	1.00 10.12	D	
	ATOM	6371		THR	159			2 170.352	1.00 17.03	D	
	ATOM	6372		THR	159			2 170.332	1.00 17.07	D	
	0	JJ , Z			100	0.190			2.00 10.07	ט	J

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	MOTA	6373	CG2		159		-70.626		1.00 17.20	D	C
	MOTA	6374	C	THR	159		-70.194		1.00 17.34	D	С
	MOTA	6375	0	THR	159		-71.409		1.00 17.77	D	0
_	MOTA	6376	N	HIS	160		-69.315		1.00 16.89	D	N
5	MOTA	6377	CA	HIS	160		-69.738		1.00 16.98	D	C
	MOTA	6378	CB	HIS	160	-10.363			1.00 16.02	D	C
	MOTA	6379	CG	HIS	160	-11.374			1.00 16.63	D	C
	MOTA	6380	CD2		160	-11.450			1.00 16.70	D	С
40	MOTA	6381	ND1		160		-69.709		1.00 15.58	D	N
10	MOTA	6382	CE1		160			164.200	1.00 16.17	D	С
	MOTA	6383		HIS	160			163.235	1.00 15.22	D	N
	MOTA	6384	C	HIS	160			165.338	1.00 16.47	D	C
	MOTA	6385	0	HIS	160			164.881	1.00 15.93	D	0
4-	MOTA	6386	N	PHE	161			164.795	1.00 16.05	D	N
15	MOTA	6387	CA	PHE	161			163.645	1.00 16.40	D	С
	ATOM	6388	СВ	PHE	161			163.125	1.00 15.95	D	С
	MOTA	6389	CG	PHE	161			162.172	1.00 15.49	D	С
	MOTA	6390		PHE	161			161.679	1.00 15.45	D	С
00	MOTA	6391		PHE	161			161.739	1.00 15.30	D	С
20	MOTA	6392		PHE	161			160.763	1.00 14.86	D	С
	ATOM	6393		PHE	161			160.826	1.00 15.33	D	С
	MOTA	6394	CZ	PHE	161			160.338	1.00 14.34	D	С
	MOTA	6395	С	PHE	161			164.066	1.00 16.58	D	С
0.5	MOTA	6396	0	PHE	161			163.377	1.00 16.53	D	0
25	MOTA	6397	N	ALA	162			165.199	1.00 16.50	D	N
	MOTA	6398	CA	ALA	162			165.687	1.00 16.70	D	C
	MOTA	6399	CB	ALA	162			167.075	1.00 15.50	D	С
	MOTA	6400	C	ALA	162			165.750	1.00 16.99	D	C
~~	MOTA	6401	0	ALA	162			165.263	1.00 16.57	D	0
30	MOTA	6402	N	ASP	163			166.329	1.00 16.55	D	N
	MOTA	6403	CA	ASP	163			166.463	1.00 17.27	D	С
	MOTA	6404	СВ	ASP	163			167.434	1.00 18.34	D	C
	MOTA	6405	CG	ASP	163			168.853	1.00 20.38	D	С
0.5	MOTA	6406		ASP	163			169.710	1.00 21.00	D	0
35	MOTA	6407		ASP	163			169.109	1.00 19.55	D	0
	MOTA	6408	C	ASP	163			165.168	1.00 16.59	D	C
	MOTA	6409	0	ASP	163			165.054	1.00 14.65	D	0
	MOTA	6410	N	ILE	164			164.196	1.00 15.61	D	N
40	ATOM	6411	CA	ILE	164			162.955	1.00 15.63	D	С
40	MOTA	6412	CB	ILE	164			162.151	1.00 14.12	D	C
	MOTA	6413	CG2		164			163.018	1.00 15.01	D	C
	MOTA	6414		ILE	164			161.666		D	C
	MOTA	6415		ILE	164			160.759	1.00 12.25	D	C
15	MOTA	6416	C	ILE	164			162.081	1.00 16.00	D	С
45	ATOM	6417	0	ILE	164			161.215	1.00 16.41	D	0
	ATOM	6418	N	ASN	165			162.300	1.00 15.80	D	N
	MOTA	6419	CA	ASN	165			. 161.542	1.00 17.21	D	C
	ATOM	6420	CB	ASN	165			161.814	1.00 16.55	D	C
EΛ	ATOM	6421	CG	ASN	165			161.068	1.00 17.71	D	C
50	MOTA	6422		ASN	165			160.122	1.00 16.40	D	0
	MOTA	6423		ASN	165			161.482	1.00 16.80	D	N
	ATOM	6424	C	ASN	165			161.990	1.00 17.61	D	С
	ATOM	6425	0	ASN	165			2 161.174	1.00 17.26	D	0
E E	MOTA	6426	N	THR	166			163.297	1.00 17.61	D	
55	ATOM	6427	CA	THR	166			7 163.818	1.00 17.94	D	C
	ATOM	6428		THR	166			7 165.355	1.00 18.82	D	C
	MOTA	6429		THR	166			7 165.748		D	_
	MOTA	6430	CG2	? THR	166	-7.015	-80.301	165.927	1.00 16.51	D	C

	MOTA	6431	C	THR	166		-79.604	_	1.00 18.00	D	C
	MOTA	6432	0	THR	166		-80.734		1.00 18.63	D	0
	MOTA	6433	N	PHE	167		-78.930		1.00 17.77	D	N
_	MOTA	6434	CA	PHE	167	-10.569			1.00 18.33	D	С
5	MOTA	6435	CB	PHE	167	-11.700			1.00 18.78	D	С
	MOTA	6436	CG	PHE	167	-12.945			1.00 19.22	D	С
	MOTA	6437	CD1		167	-13.727			1.00 19.14	D	С
	ATOM	6438	CD2		167	-13.343			1.00 19.37	D	С
4.0	ATOM	6439	CE1		167	-14.892			1.00 18.14	D	С
10	MOTA	6440			167	-14.499			1.00 19.96	D	C
	MOTA	6441	CZ	PHE	167	-15.279			1.00 20.10	D	C
	MOTA	6442	С	PHE	167			161.550	1.00 18.30	D	C
	MOTA	6443	0	PHE	167			161.211	1.00 18.49	D	0
	MOTA	6444	N	MET	168			160.694	1.00 17.27	D	N
15	MOTA	6445	CA	MET	168			159.269	1.00 17.47	D	С
	MOTA	6446	CB	MET	168			158.484	1.00 17.11	D	С
	MOTA	6447	CG	MET	168			158.163	1.00 17.30	D	С
	MOTA	6448	SD	MET	168			157.061	1.00 15.92	D	S
	MOTA	6449	CE	MET	168			158.216	1.00 15.99	D	С
20	MOTA	6450	С	MET	168			158.949	1.00 16.51	D	С
	MOTA	6451	0	MET	168			158.120	1.00 15.80	D	0
	ATOM	6452	N	VAL	169			159.582	1.00 16.26	D	N
	MOTA	6453	CA	VAL	169			159.313	1.00 17.34	D	С
	MOTA	6454	CB	VAL	169			160.042	1.00 17.83	D	С
25	MOTA	6455		VAL	169			159.813	1.00 18.06	D	С
	MOTA	6456		VAL	169			159.527	1.00 17.43	D	C
	MOTA	6457	С	VAL	169			159.739	1.00 17.64	D	C
	MOTA	6458	0	VAL	169			159.076	1.00 16.56	D	0
~~	MOTA	6459	N	LEU	170			160.847	1.00 17.79	D	N
30	MOTA	6460	CA	LEU	170			161.302	1.00 19.01	D	C
	MOTA	6461	CB	LEU	170			162.656	1.00 20.40	D	C
	MOTA	6462	CG	LEU	170			163.770	1.00 22.26	D	С
	ATOM	6463		LEU	170			165.038	1.00 23.26	D	С
٥.5	MOTA	6464		LEU	170			164.010	1.00 23.21	D	С
35	MOTA	6465	C	LEU	170			160.258	1.00 19.01	D	С
	ATOM	6466	0	LEU	170			160.021	1.00 19.62	D	0
	ATOM	6467	N	GLN	171			159.621	1.00 18.39	D	N
	ATOM	6468	CA	GLN	171			158.591	1.00 19.08	D	C
40	MOTA	6469	CB	GLN	171			158.225	1.00 19.14	D	C
40	MOTA	6470	CG	GLN	171			159.342	1.00 20.31	D	C
	ATOM	6471	CD	GLN	171			159.744	1.00 20.59	D	С
	ATOM	6472		GLN	171			158.960		D	_
	ATOM	6473		GLN	171			160.955	1.00 19.36	D	N
45	ATOM	6474	C	GLN	171			157.346	1.00 19.36	D	C
45	ATOM	6475	0	GLN	171			156.736	1.00 18.90	D	0
	ATOM	6476	N	VAL	172			156.968	1.00 18.78	D	N
	ATOM	6477	CA	VAL	172			155.817	1.00 19.25	D	C
	ATOM	6478	CB	VAL	172			155.559	1.00 19.23	D	C
50	ATOM	6479		VAL	172			2 154.527	1.00 17.92	D	C
50	MOTA	6480		VAL	172			155.063	1.00 18.09	D	C
	MOTA	6481	C	VAL	172			156.097	1.00 19.25	D	C
	MOTA	6482	0	VAL	172			155.213	1.00 17.74	D	0
	MOTA	6483	N	ILE	173			157.333	1.00 19.42	D	N
EE	MOTA	6484	CA	ILE	173			157.748		D	C
55	ATOM	6485	CB	ILE	173			159.227		D	C
	MOTA	6486		ILE	173			159.756		D	C
	ATOM	6487		LILE	173			5 159.332		D	C
	ATOM	6488	CD1	LILE	173	-5.807	-87.079	160.744	1.00 20.26	D	C

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	MOTA	6489	С	ILE	173		-88.727		1.00 21.22	D	С
	MOTA	6490	0	ILE	173		-89.847		1.00 21.19	D	0
	MOTA	6491	N	LYS	174	-10.557	-88.211	157.982	1.00 21.41	D	N
	MOTA	6492	CA	LYS	174	-11.808	-88.946	157.835	1.00 22.16	D	С
5	MOTA	6493	CB	LYS	174	-12.933	-88.231	158.599	1.00 23.61	D	С
	ATOM	6494	CG	LYS	174	-12.720	-88.228	160.119	1.00 25.13	D	C
	MOTA	6495	CD	LYS	174	-13.770	-87.422	160.870	1.00 26.92	D	С
	ATOM	6496	CE	LYS	174	-15.122	-88.112	160.854	1.00 30.21	D	С
	MOTA	6497	NZ	LYS	174	-16.122	-87.399	161.705	1.00 31.50	D	N
10	MOTA	6498	С	LYS	174	-12.156	-89.082	156.348	1.00 22.88	D	С
. •	MOTA	6499	ō	LYS	174	-12.806			1.00 21.83	D	0
	ATOM	6500	N	PHE	175	-11.715			1.00 23.02	D	N
	ATOM	6501	CA	PHE	175			154.107	1.00 23.18	D	С
	ATOM	6502	СВ	PHE	175			153.498	1.00 22.95	D	С
15	ATOM	6503	CG	PHE	175			152.024	1.00 22.60	D	С
10	ATOM	6504	CD1		175			151.086	1.00 21.67	D	Ċ
	MOTA	6505		PHE	175			151.575	1.00 22.02	D	Ċ
	ATOM	6506		PHE	175			149.719	1.00 22.60	D	c
	ATOM	6507	CE2	PHE	175			150.215	1.00 22.00	D	C
20		6508	CZ	PHE	175			149.284	1.00 22.15	D	C
20	ATOM							153.479	1.00 22.43	D	C
	MOTA	6509 6510	C	PHE	175 175			152.725	1.00 23.77	D	0
	MOTA		0	PHE				152.725	1.00 23.33		N
	ATOM	6511	N	THR	176				1.00 24.37	D	C
05	MOTA	6512	CA	THR	176			153.248		D	
25	MOTA	6513	CB	THR	176			153.542	1.00 25.42	D	C
	ATOM	6514	OG1		176			154.956	1.00 26.76	D	0
	ATOM	6515	CG2		176			152.878	1.00 25.58	D	C
	ATOM	6516	C	THR	176			153.733	1.00 26.23	D	С
~~	MOTA	6517	0	THR	176			153.012	1.00 26.27	D	0
30	MOTA	6518	N	LYS	177			154.935	1.00 26.44	D	N
	MOTA	6519	CA	LYS	177			155.453	1.00 27.74	D	С
	ATOM	6520	CB	LYS	177			156.867	1.00 27.72	D	C
	MOTA	6521	CG	LYS	177			157.354	1.00 28.98	D	C
	MOTA	6522	CD	LYS	177			158.723	1.00 28.59	D	C
35	MOTA	6523	CE	LYS	177	-12.497	-95.914	159.106	1.00 28.79	D	С
	MOTA	6524	NZ	LYS	177			158.341	1.00 27.25	D	N
	ATOM	6525	С	LYS	177	-11.287	-93.906	154.546	1.00 27.56	D	С
	MOTA	6526	0	LYS	177	-11.251	-95.124	154.381	1.00 27.25	D	0
	MOTA	6527	N	ASP	178	-12.183	-93.111	. 153.966	1.00 27.88	D	N
40	ATOM	6528	CA	ASP	178	-13.218	-93.616	153.069	1.00 28.74	D	C
	ATOM	6529	СВ	ASP	178	-14.388	-92.627	153.006	1.00 29.09	D	С
	ATOM	6530	CG	ASP	178	-15.247	-92.647	154.258	1.00 30.01	D	C
	ATOM	6531	OD1	ASP	178	-14.983	-93.459	155.168	1.00 30.64	D	0
	MOTA	6532		ASP	178			154.327	1.00 30.67	D	0
45	ATOM	6533	С	ASP	178			151.644	1.00 28.80	D	С
. •	ATOM	6534	0	ASP	178	-13.556	-94.076	5 150.743	1.00 29.46	D	0
	ATOM	6535	N	LEU	179			151.434	1.00 28.53	D	N
	ATOM	6536	CA	LEU	179			5 150.112	1.00 27.98	D	С
	MOTA	6537	СВ	LEU	179			2 149.696		D	C
50	ATOM	6538	CG	LEU	179			5 149.737		D	Ċ
50	ATOM	6539		LEU	179			3 149.145			c
		6540		LEU	179			5 143.143 5 148.959		D	c
	ATOM	6541	CDZ	LEU	179			3 140.939 3 150.185			C
	ATOM	6542	0	LEU	179			2 150.183			0
55	MOTA				180			z 130.683 5 149.682			
၁၁	ATOM	6543		PRO							
	ATOM	6544		PRO	180			7 148.999 7 140.601			
	ATOM	6545		PRO	180			7 149.691			
	ATOM	6546	CB	PRO	180	-TI.085	, -98.71	3 148.793	1.00 29.68	D	С

	ATOM	6547	CG	PRO	180	-12.420 -98.052 149.056 1.00 28.81 D	С
	ATOM	6548	C	PRO	180	-8.691 -97.991 149.223 1.00 29.88 D	Ċ
	MOTA	6549	ō	PRO	180	-7.838 -98.543 149.918 1.00 29.69 p	Ö
	ATOM	6550	N	VAL	181	-8.399 -97.429 148.055 1.00 30.49 D	N
5	ATOM	6551	CA	VAL	181	-7.036 -97.489 147.554 1.00 31.05 D	C
-	ATOM	6552	СВ	VAL	181	-6.943 -96.959 146.105 1.00 32.32 D	Č
	ATOM	6553	CG1		181	-7.329 -95.478 146.032 1.00 33.43 D	č
	ATOM	6554		VAL	181	-5.545 -97.204 145.582 1.00 33.81 D	Ċ
	ATOM	6555	C	VAL	181	-6.030 -96.777 148.462 1.00 30.44 D	C
10	ATOM	6556	Ö	VAL	181	-4.866 -97.172 148.533 1.00 31.06 D	Ö
. •	ATOM	6557	N	PHE	182		
	MOTA	6558	CA	PHE	182		N C
	ATOM	6559	СВ	PHE	182	A A A A A A A A A A A A A A A A A A A	C
	ATOM	6560	CG	PHE	182		C
15	MOTA	6561		PHE	182		
10	MOTA	6562		PHE	182		C
	ATOM	6563		PHE	182		C
	ATOM	6564	CE2	PHE	182		С
	MOTA	6565	CZ	PHE	182		С
20						-3.942 -91.599 153.558 1.00 24.75 D	С
20	MOTA	6566	С	PHE	182	-5.347 -96.000 151.273 1.00 27.26 D	С
	MOTA	6567	0	PHE	182	-4.218 -96.201 151.707 1.00 25.52 D	0
	ATOM	6568	N	ARG	183	-6.426 -96.599 151.772 1.00 27.61 D	N
	ATOM	6569	CA	ARG	183	-6.348 -97.535 152.900 1.00 29.82 D	С
25	MOTA	6570	CB	ARG	183	-7.751 -97.955 153.342 1.00 29.81 D	С
25	MOTA	6571	CG	ARG	183	-8.610 -96.819 153.834 1.00 30.04 D	С
	MOTA	6572	CD	ARG	183	-8.065 -96.222 155.126 1.00 31.16 D	С
	ATOM	6573	NE	ARG	183	-8.007 -97.174 156.241 1.00 32.26 D	N
	MOTA	6574	CZ	ARG	183	-9.048 -97.832 156.752 1.00 33.84 D	С
20	ATOM	6575		ARG	183	-10.271 -97.670 156.256 1.00 34.26 D	N
30	MOTA	6576	NH2		183	-8.870 -98.641 157.788 1.00 32.25 D	N
	ATOM	6577	C	ARG	183	-5.538 -98.791 152.611 1.00 30.35 D	-
	MOTA	6578	0	ARG	183	-5.025 -99.428 153.536 1.00 30.29 D	-
	ATOM	6579	N	SER	184	-5.436 -99.155 151.335 1.00 30.53 D	
0.5	MOTA	6580	CA	SER	184	-4.680-100.341 150.937 1.00 31.20 D	
35	ATOM	6581	CB	SER	184	-4.960-100.687 149.472 1.00 31.90 D	C
	ATOM	6582	OG	SER	184	-6.311-101.079 149.299 1.00 34.38 D	_
	MOTA	6583	С	SER	184	-3.187-100.149 151.129 1.00 31.02 D	
	ATOM	6584	0	SER	184	-2.440-101.119 151.218 1.00 31.50 D	0
	MOTA	6585	N	LEU	185	-2.752 -98.894 151.185 1.00 30.38 D	N
40	MOTA	6586	CA	LEU	185	-1.340 -98.580 151.369 1.00 30.35 D	C
	MOTA	6587	CB	LEU	185	-1.076 -97.105 151.031 1.00 30.07 D	C
	ATOM	6588	CG	LEU	185	-1.534 -96.535 149.683 1.00 30.04 D	C
	MOTA	6589		LEU	185	-1.345 -95.013 149.690 1.00 28.56 D	C
	MOTA	6590	CD2	LEU	185	-0.748 -97.180 148.546 1.00 28.92 D	C
45	MOTA	6591	С	LEU	185	-0.974 -98.817 152.831 1.00 30.64 D	C
	MOTA	6592	0	LEU	185	-1.848 -98.832 153.696 1.00 30.76 D	
	ATOM	6593	N	PRO	186	0.320 -99.018 153.125 1.00 31.27 D	
	MOTA	6594	CD	PRO	186	1.483 -99.091 152.224 1.00 31.66 D	
	ATOM	6595	CA	PRO	186	0.724 -99.235 154.515 1.00 31.60 D	
50	MOTA	6596	CB	PRO	186	2.220 -99.522 154.401 1.00 31.16 D	
	MOTA	6597	CG	PRO	186	2.620 -98.801 153.164 1.00 32.29 D	
	MOTA	6598	С	PRO	186	0.419 -97.956 155.299 1.00 31.66 D	
	ATOM	6599	0	PRO	186	0.609 -96.855 154.791 1.00 31.55 D	
	ATOM	6600	N	ILE	187	-0.054 -98.106 156.531 1.00 31.45 D	
55	ATOM	6601	CA	ILE	187	-0.425 -96.959 157.351 1.00 31.05 D	
	ATOM	6602	CB	ILE	187	-0.893 -97.417 158.766 1.00 29.97 D	
	ATOM	6603		ILE	187	0.292 -97.909 159.591 1.00 29.23 D	
	ATOM	6604		ILE	187	-1.606 -96.264 159.476 1.00 28.40 D	
					<b>~</b> • •	JU.204 1JJ.4/U 1.00 20.4U D	

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	ATOM	6605	CD1	TTE	187	_2 392	-96.690	160 706	1.00 27.64	D	С
	ATOM	6606	CDI	ILE	187		-95.861		1.00 27.84	D	c
	ATOM	6607	0	ILE	187		-94.683		1.00 31.30	D	o
							-96.220		1.00 31.41		
5	ATOM	6608	N	GLU	188		-95.220		1.00 32.22	D	N
3	ATOM	6609	CA	GLU	188					D	C
	MOTA	6610	CB	GLU	188		-95.847		1.00 35.42	D	C
	ATOM	6611	CG	GLU	188		-96.637		1.00 38.74	D	C
	ATOM	6612	CD	GLU	188		-97.046		1.00 41.52	D	C
	MOTA	6613	OE1		188		-96.137		1.00 42.54	D	0
10	MOTA	6614	OE2	GLU	188		-98.267		1.00 42.40	D	0
	MOTA	6615	С	GLU	188		-94.324		1.00 33.32	D	C
	MOTA	6616	0	GLU	188	3.398	-93.158	156.381	1.00 33.14	D	0
	MOTA	6617	N	ASP	189	2.638	-94.888	155.163	1.00 32.68	D	N
	MOTA	6618	CA	ASP	189	2.596	-94.130	153.921	1.00 32.83	D	C
15	MOTA	6619	CB	ASP	189	2.505	-95.072	152.713	1.00 35.38	D	С
	ATOM	6620	CG	ASP	189	3.800	-95.828	152.465	1.00 37.57	D	С
	MOTA	6621		ASP	189		-96.542		1.00 40.86	D	0
	ATOM	6622		ASP	189		-95.705		1.00 37.64	D	0
	ATOM	6623	C	ASP	189		-93.238		1.00 30.96	D	Ċ
20	ATOM	6624	Ö	ASP	189		-92.078		1.00 30.94	D	ō
	ATOM	6625	N	GLN	190		-93.786		1.00 29.38	D	N
	MOTA	6626	CA	GLN	190		-93.024		1.00 28.01	D	C
	MOTA	6627	CB	GLN	190			155.350	1.00 28.17	D	C
	MOTA	6628	CG	GLN	190			154.661	1.00 28.90	D	C
25	ATOM	6629	CD	GLN	190			155.323	1.00 28.30		C
23										D	
	MOTA	6630		GLN	190			155.094	1.00 29.27	D	0
	MOTA	6631	NE2		190			156.130	1.00 28.27	D	И
	MOTA	6632	C	GLN	190			155.447	1.00 27.34	D	C
00	MOTA	6633	0	GLN	190			155.110	1.00 25.56	D	0
30	ATOM	6634	N	ILE	191			156.523	1.00 26.72	D	N
	MOTA	6635	CA	ILE	191			157.389	1.00 26.98	D	C
	ATOM	6636	CB	ILE	191			158.693	1.00 27.03	D	С
	MOTA	6637	CG2		191			159.487	1.00 26.61	D	С
C_	MOTA	6638	CG1	ILE	191			159.520	1.00 26.71	D	C
35	MOTA	6639	CD1	ILE	191			160.693	1.00 27.05	D	C
	ATOM	6640	С	ILE	191	1.396	-89.835	156.690	1.00 26.93	D	С
	ATOM	6641	0	ILE	191			156.859	1.00 25.97	D	0
	MOTA	6642	N	SER	192	2.343	-90.348	155.907	1.00 25.82	D	N
	ATOM	6643	CA	SER	192	3.270	-89.486	155.183	1.00 26.80	D	C
40	MOTA	6644	CB	SER	192	4.338	-90.325	154.473	1.00 27.15	D	C
	ATOM	6645	OG	SER	192	5.203	-90.939	155.415	1.00 29.76	D	0
	ATOM	6646	С	SER	192	2.531	-88.623	154.154	1.00 25.97	D	C
	MOTA	6647	0	SER	192	2.788	-87.425	154.037	1.00 25.52	D	0
	MOTA	6648	N	LEU	193	1.621	-89.238	153.407	1.00 24.97	D	N
45	MOTA	6649	CA	LEU	193	0.857		152.402	1.00 24.60	D	C
	ATOM	6650	CB	LEU	193			151.539	1.00 24.47	D	C
	ATOM	6651	CG	LEU	193			150.667	1.00 25.09	D	Č
	MOTA	6652		LEU	193			149.936	1.00 21.73	D	c
	ATOM	6653		LEU	193			149.673	1.00 24.35	D	c
50	ATOM	6654	C	LEU	193			153.032	1.00 24.39	D	c
50		6655	Ö	LEU	193			152.476		D	
	ATOM				193				1.00 23.01 1.00 24.06		0
	ATOM	6656	N	LEU				154.181		D	N
	ATOM	6657	CA	LEU	194			154.862	1.00 24.67	D	C
EE	ATOM	6658	CB	LEU	194			156.014		D	C
55	ATOM	6659	CG	LEU	194			155.880		D	C
	MOTA	6660		LEU	194			156.161		D	C
	MOTA	6661		LEU	194			154.492		D	-
	ATOM	6662	С	LEU	194	-0.766	-85.711	155.375	1.00 24.96	D	C

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	MOTA	6663	0	LEU	194		-84.559		1.00 24.50	D	0
	MOTA	6664	N	LYS	195		-85.999		1.00 24.56	D	N
	MOTA	6665	CA	LYS	195		-84.971		1.00 25.60	D	С
_	MOTA	6666	CB	LYS	195		-85.642		1.00 26.77	D	С
5	MOTA	6667	CG	LYS	195		-84.704		1.00 31.05	D	С
	MOTA	6668	CD	LYS	195		-85.476		1.00 33.47	D	С
	MOTA	6669	CE	LYS	195		-86.190		1.00 34.73	D	С
	MOTA	6670	NZ	LYS	195			158.448	1.00 37.37	D	N
40	MOTA	6671	С	LYS	195			155.349	1.00 24.73	D	С
10	MOTA	6672	0	LYS	195			155.569	1.00 23.70	D	0
	MOTA	6673	N	GLY	196			154.152	1.00 22.84	D	N
	ATOM	6674	CA	GLY	196			153.062	1.00 21.25	D	С
	MOTA	6675	С	GLY	196			152.282	1.00 20.83	D	С
. –	MOTA	6676	0	GLY	196			151.705	1.00 20.29	D	0
15	MOTA	6677	N	ALA	197			152.277	1.00 20.54	D	N
	MOTA	6678	CA	ALA	197			151.504	1.00 19.62	D	C
	MOTA	6679	CB	ALA	197			150.522	1.00 18.07	D	С
	MOTA	6680	С	ALA	197			152.248	1.00 18.74	D	С
	MOTA	6681	0	ALA	197			151.634	1.00 17.93	D	0
20	MOTA	6682	N	ALA	198			153.542	1.00 18.19	D	N
	MOTA	6683	CA	ALA	198			154.311	1.00 17.94	D	C
	MOTA	6684	CB	ALA	198			155.805	1.00 18.85	D	C
	MOTA	6685	C	ALA	198	-3.767	-80.588	154.102	1.00 16.59	D	С
	ATOM	6686	0	ALA	198			153.758	1.00 15.86	D	0
25	MOTA	6687	N	VAL	199			154.320	1.00 16.08	D	N
	ATOM	6688	CA	VAL	199	-2.951	-78.306	154.142	1.00 16.51	D	C
	ATOM	6689	CB	VAL	199	-1.648	-77.545	154.467	1.00 16.75	D	С
	MOTA	6690	CG1	VAL	199	-1.814	-76.066	154.138	1.00 18.24	D	C
	ATOM	6691	CG2	VAL	199			155.945	1.00 18.67	D	C
30	MOTA	6692	С	VAL	199	-3.395	-77.971	152.709	1.00 16.12	D	C
	MOTA	6693	0	VAL	199	-4.300	-77.156	152.498	1.00 14.02	D	0
	ATOM	6694	N	GLU	200	-2.755	-78.601	151.727	1.00 15.48	D	N
	ATOM	6695	CA	GLU	200	-3.097	-78.366	150.330	1.00 16.73	D	C
	MOTA	6696	СВ	GLU	200	-2.150	-79.157	149.417	1.00 16.50	D	C
35	MOTA	6697	CG	GLU	200			149.252	1.00 17.80	D	C
	MOTA	6698	CD	GLU	200			148.533	1.00 19.22	D	C
	ATOM	6699	OE1	GLU	200	-0.122	-80.054	147.521	1.00 20.12	D	0
	ATOM	6700	OE2	GLU	200	1.404	-79.481	148.975	1.00 18.83	D	0
	MOTA	6701	C	GLU	200	-4.557	-78.721	150.035	1.00 15.98	D	С
40	MOTA	6702	0	GLU	200	-5.285	-77.941	149.402	1.00 15.85	D	0
	MOTA	6703	N	ILE	201	-4.989	-79.890	150.499	1.00 15.63	D	N
	ATOM	6704	CA	ILE	201	-6.364	-80.340	150.293	1.00 15.51	D	С
	MOTA	6705	CB	ILE	201	-6.575	-81.743	150.927	1.00 15.91	D	C
	ATOM	6706	CG2	ILE	201	-8.058	-82.111	. 150.932	1.00 15.40	D	C
45	ATOM	6707	CG1	. ILE	201	-5.766	-82.783	150.129	1.00 18.12	D	C
	ATOM	6708	CD1	ILE	201	-5.472	-84.091	150.870	1.00 15.80	D	С
	ATOM	6709	C	ILE	201	-7.342	-79.327	150.895	1.00 15.21	D	С
	MOTA	6710	0	ILE	201	-8.321	-78.927	7 150.251	1.00 14.04	D	0
	ATOM	6711	N	CYS	202	-7.065	-78.892	2 152.119	1.00 14.29	D	N
50	ATOM	6712	CA	CYS	202	-7.934	-77.927	7 152.776	1.00 15.16	D	С
	ATOM	6713	CB	CYS	202	-7.383	-77.554	154.155	1.00 14.22	D	С
	ATOM	6714	SG	CYS	202			2 155.348	1.00 16.23	D	S
	MOTA	6715	С	CYS	202	-8.124	-76.679	151.931	1.00 14.68	D	
	ATOM	6716	0	CYS	202			151.809	1.00 13.66	D	
55	ATOM	6717	N	HIS	203			3 151.343	1.00 14.61	D	
	ATOM	6718	CA	HIS	203			1 150.509		D	
	ATOM	6719		HIS	203			5 150.140		D	
	ATOM	6720		HIS	203			3 151.227		D	
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	MOTA	6721	CD2 HIS	203		-73.810		1.00 17.06		С
	MOTA	6722	ND1 HIS	203		-72.389		1.00 17.22		N
	MOTA	6723	CE1 HIS	203		-71.905		1.00 17.87	D	С
_	ATOM	6724	NE2 HIS	203		-72.749		1.00 18.68	D	N
5	ATOM	6725	C HIS	203		-75.249		1.00 15.18	D	С
	MOTA	6726	O HIS	203		-74.395		1.00 14.74	D	0
	MOTA	6727	N ILE	204		-76.441		1.00 15.83	D	N
	MOTA	6728	CA ILE	204		-76.797		1.00 15.89	D	С
	ATOM	6729	CB ILE	204		-78.195		1.00 16.42	D	С
10	ATOM	6730	CG2 ILE	204			145.904	1.00 15.76	D	С
	MOTA	6731	CG1 ILE	204			146.458	1.00 16.43	D	С
	MOTA	6732	CD1 ILE	204		-79.496		1.00 18.76	D	С
	MOTA	6733	C ILE	204	-10.108			1.00 16.28	D	C
	MOTA	6734	O ILE	204	-10.943			1.00 15.45	D	0
15	MOTA	6735	N VAL	205	-10.423			1.00 15.70	D	N
	MOTA	6736	CA VAL	205	-11.799			1.00 16.11	D	C
	MOTA	6737	CB VAL	205	-11.895			1.00 16.88	D	С
	MOTA	6738	CG1 VAL	205	-13.274			1.00 16.36	D	C
	MOTA	6739	CG2 VAL	205	-11.661			1.00 16.38	D	С
20	MOTA	6740	C VAL	205	-12.340	-76.103	149.906	1.00 16.68	D	C
	MOTA	6741	O VAL	205	-13.452			1.00 16.11	D	0
	MOTA	6742	N LEU	206	-11.544			1.00 16.97	D	N
	MOTA	6743	CA LEU	206	-11.962			1.00 18.74	D	C
	MOTA	6744	CB LEU	206	-10.932			1.00 19.59	D	C
25	MOTA	6745	CG LEU	206	-11.376			1.00 21.97	D	С
	MOTA	6746	CD1 LEU	206			153.922	1.00 23.35	D	С
	MOTA	6747	CD2 LEU	206			154.403	1.00 22.61	D	С
	MOTA	6748	C LEU	206			149.886	1.00 18.61	D	C
	MOTA	6749	O LEU	206			150.023	1.00 18.78	D	0
30	MOTA	6750	N ASN	207			148.740	1.00 17.84	D	N
	MOTA	6751	CA ASN	207			147.569	1.00 18.81	D	С
	MOTA	6752	CB ASN	207			146.358	1.00 16.45	D	С
	MOTA	6753	CG ASN	207			145.167	1.00 17.11	D	С
	MOTA	6754	OD1 ASN	207			144.162	1.00 13.45	D	0
35	ATOM	6755	ND2 ASN	207			145.283	1.00 15.61	D	N
	MOTA	6756	C ASN	207			147.206	1.00 19.76	D	С
	MOTA	6757	O ASN	207			146.727	1.00 19.75	D	0
	MOTA	6758	N THR	208			147.442	1.00 20.34	D	Ŋ
	MOTA	6759	CA THR	208			147.111	1.00 22.21	D	C
40	MOTA	6760	CB THR	208			147.240	1.00 23.45	D	С
	MOTA	6761	OG1 THR	208			148.599	1.00 23.83	D	0
	MOTA	6762	CG2 THR	208			146.332	1.00 24.88	D	
	MOTA	6763	C THR	208			147.938	1.00 21.89	D	С
4-	MOTA	6764	O THR	208			147.626	1.00 23.81	D	0
45	MOTA	6765	N THR	209			148.981	1.00 20.43	D	N
	MOTA	6766	CA THR	209			149.799	1.00 19.96	D	С
	MOTA	6767	CB THR	209			151.320	1.00 18.96	D	C
	MOTA	6768	OG1 THR	209			151.632	1.00 17.92	D	0
	MOTA	6769	CG2 THR	209			151.732	1.00 18.80	D	C
50	MOTA	6770	C THR	209			149.455	1.00 19.40	D	С
	MOTA	6771	O THR	209			150.008		D	0
	MOTA	6772	N PHE	210			148.544	1.00 19.55	D	N
	ATOM	6773		210			148.151		D	С
	ATOM	6774		210			147.433		D	С
55	MOTA	6775		210			147.289		D	С
	MOTA	6776		210			148.396		D	C
	MOTA	6777		210			146.046		D	C
	MOTA	6778	CE1 PHE	210	-10.054	-65.350	148.265	1.00 17.81	D	C

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	MOTA	6779	CE2	PHE	210	-10.500	-65.281	145.902	1.00 17.22	D	С
	MOTA	6780	CZ	PHE	210	-9.841	-64.775	147.011	1.00 18.25	D	C
	MOTA	6781	С	PHE	210	-14.722	-66.958	147.240	1.00 21.26	D	C
	MOTA	6782	0	PHE	210	-15.199	-67.428	146.212	1.00 20.72	D	0
<b>5</b> .	MOTA	6783	N	CYS	211	-14.926	-65.708	147.630	1.00 23.14	D	N
	ATOM	6784	CA	CYS	211	-15.736	-64.776	146.862	1.00 25.64	D	С
	MOTA	6785	CB	CYS	211	-16.594	-63.928	147.803	1.00 26.00	D	С
	MOTA	6786	SG	CYS	211	-17.553	-62.630	146.984	1.00 28.41	D	S
	MOTA	6787	С	CYS	211	-14.812	-63.880	146.044	1.00 27.00	D	С
10	ATOM	6788	0	CYS	211	-14.064	-63.070	146.592	1.00 26.49	D	0
	ATOM	6789	N	LEU	212	-14.865	-64.037	144.728	1.00 28.64	D	N
	MOTA	6790	CA	LEU	212	-14.032	-63.250	143.830	1.00 30.45	D	С
	MOTA	6791	CB	LEU	212	-14.267	-63.683	142.383	1.00 30.47	D	С
	MOTA	6792	CG	LEU	212	-13.780	-65.084	142.006	1.00 30.70	D	С
15	MOTA	6793	CD1	LEU	212	-14.109	-65.364	140.550	1.00 31.20	D	С
	ATOM	6794	CD2	LEU	212	-12.285	-65.183	142.237	1.00 30.24	D	С
	MOTA	6795	С	LEU	212	-14.264	-61.748	143.943	1.00 31.68	D	С
	MOTA	6796	0	LEU	212	-13.327	-60.964	143.793	1.00 31.74	D	0
	ATOM	6797	N	GLN	213	-15.505	-61.349	144.216	1.00 32.60	D	N
20	MOTA	6798	CA	GLN	213	-15.844	-59.935	144.315	1.00 33.45	D	С
	ATOM	6799	CB	GLN	213	-17.366	-59.763	144.407	1.00 36.26	D	С
	MOTA	6800	CG	GLN	213	-17.835	-58.346	144.099	1.00 40.79	D	С
	ATOM	6801	CD	GLN	213	-19.349	-58.213	144.055	1.00 43.35	D	C
	MOTA	6802	OE1	GLN	213	-20.027	-58.316	145.082	1.00 44.82	D	0
25	MOTA	6803	NE2	GLN	213	-19.889	-57.984	142.859	1.00 44.19	D	N
	MOTA	6804	С	GLN	213	-15.175	-59.209	145.478	1.00 32.04	D	C
	ATOM	6805	0	GLN	213	-14.750	-58.067	145.331	1.00 31.86	D	0
	ATOM	6806	N	THR	214	-15.076	-59.867	146.629	1.00 30.21	D	N
	ATOM	6807	CA	THR	214	-14.467	-59.249	147.803	1.00 28.64	D	С
30	ATOM	6808	CB	THR	214	-15.435	-59.283	149.002	1.00 28.97	D	C
	ATOM	6809	OG1	THR	214	-15.878	-60.630	149.218	1.00 28.69	D	0
	ATOM	6810	CG2	THR	214	-16.646	-58.390	148.736	1.00 28.63	D	C
	ATOM	6811	С	THR	214	-13.146	-59.870	148.253	1.00 27.89	D	С
	MOTA	6812	0	THR	214	-12.523	-59.371	149.184	1.00 27.24	D	0
35	MOTA	6813	N	GLN	215	-12.719	-60.949	147.600	1.00 27.59	D	N
	MOTA	6814	CA	GLN	215	-11.475	-61.627	147.969	1.00 27.84	D	С
	MOTA	6815	CB	GLN	215	-10.285	-60.671	147.836	1.00 29.73	D	С
	MOTA	6816	CG	GLN	215	-10.102	-60.059	146.458	1.00 32.60	D	C
	ATOM	6817	CD	GLN	215	-9.768	-61.086	145.407	1.00 34.25	D	С
40	ATOM	6818	OE1	GLN	215	-10.590	-61.407	144.550	1.00 36.19	D	0
	MOTA	6819	NE2	GLN	215	-8.555	-61.617	145.471	1.00 35.54	D	N
	MOTA	6820	С	GLN	215			. 149.419	1.00 26.63	D	C
	MOTA	6821	0	GLN	215	-10.579	-62.068	150.158	1.00 25.86	D	0
	ATOM	6822	N	ASN	216	-12.734	-62.600	149.810	1.00 25.65	D	N
45	MOTA	6823	CA	ASN	216	-12.985	-63.097	151.160	1.00 25.70	D	C
	ATOM	6824	CB	ASN	216	-14.110	-62.292	2 151.812	1.00 27.21	D	С
	MOTA	6825	CG	ASN	216	-13.691	-60.904	152.208	1.00 29.76	D	C
	MOTA	6826	OD1	ASN	216	-14.536	-60.054	152.492	1.00 31.19	D	0
	ATOM	6827	ND2	ASN	216	-12.384	-60.663	3 152.258	1.00 31.91	Ď	N
50	MOTA	6828	C	ASN	216	-13.435	-64.550	151.133	1.00 24.52	D	C
	MOTA	6829		ASN	216	-13.870	-65.05	l 150.103	1.00 23.28	D	0
	MOTA	6830		PHE	217	-13.330	-65.218	3 152.276	1.00 23.35	D	N
	MOTA	6831		PHE	217	-13.799	-66.593	152.396	1.00 22.98	D	С
	MOTA	6832	CB	PHE	217	-12.782	-67.47	5 153.131	1.00 20.37	D	С
55	MOTA	6833	CG	PHE	217	-11.514	-67.72	7 152.352	1.00 19.81	D	C
	MOTA	6834		PHE	217			152.396		D	С
	MOTA	6835		PHE	217			9 151.556		D	С
	MOTA	6836		L PHE	217			3 151.655		D	

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	MOTA	6837	CE2		217	-10.228			1.00 19.86	D	С
	MOTA	6838	CZ	PHE	217		-68.176		1.00 19.40	D	С
	MOTA	6839	С	PHE	217	-15.094			1.00 23.95	D	С
_	MOTA	6840	0	PHE	217	-15.074			1.00 23.31	D	0
5	ATOM	6841	N	LEU	218	-16.216			1.00 24.22	D	N
	MOTA	6842	CA	LEU	218	-17.508			1.00 25.10	D	С
	MOTA	6843	CB	LEU	218	-18.586			1.00 26.14	D	С
	MOTA	6844	CG	LEU	218		-64.972		1.00 27.28	D	С
	MOTA	6845	CD1		218		-64.588		1.00 26.98	D	С
10	MOTA	6846	CD2		218		-63.883		1.00 27.40	D	С
	MOTA	6847	С	LEU	218			153.786	1.00 25.53	D	С
	MOTA	6848	0	LEU	218		-69.087		1.00 25.80	D	0
	MOTA	6849	N	CYS	219			155.103	1.00 25.26	D	N
	MOTA	6850	CA	CYS	219			155.733	1.00 25.79	D	С
15	ATOM	6851	CB	CYS	219			156.477	1.00 25.11	D	C
	MOTA	6852	SG	CYS	219			155.391	1.00 24.90	D	S
	MOTA	6853	С	CYS	219			156.674	1.00 25.79	D	С
	MOTA	6854	0	CYS	219	-19.103	-69.165	157.857	1.00 25.32	D	0
	MOTA	6855	N	GLY	220			156.131	1.00 25.40	D	N
20	MOTA	6856	CA	GLY	220	-21.636	-69.660	156.922	1.00 24.69	D	С
	MOTA	6857	С	GLY	220	-21.761	-68.211	157.341	1.00 24.74	D	С
	ATOM	6858	0	GLY	220			156.492	1.00 24.53	D	0
	ATOM	6859	N	PRO	221	-21.795	-67.933	158.649	1.00 24.44	D	N
	ATOM	6860	CD	PRO	221	-21.952	-68.897	159.758	1.00 24.16	D	С
25	ATOM	6861	CA	PRO	221	-21.912	-66.552	159.119	1.00 24.43	D	С
	MOTA	6862	CB	PRO	221	-22.545	-66.722	160.495	1.00 25.03	D	С
	MOTA	6863	CG	PRO	221	-21.911	-68.011	160.983	1.00 24.72	D	C
	ATOM	6864	С	PRO	221	-20.573	-65.816	159.192	1.00 24.14	D	C
	MOTA	6865	0	PRO	221			159.401	1.00 24.82	D	0
30	ATOM	6866	N	LEU	222	-19.482	-66.552	159.014	1.00 23.02	D	N
	ATOM	6867	CA	LEU	222	-18.143	-65.975	159.097	1.00 22.19	D	C
	MOTA	6868	CB	LEU	222	-17.171	-67.006	159.675	1.00 20.44	D	C
	ATOM	6869	CG	LEU	222	-17.611	-67.653	160.991	1.00 19.18	D	C
	ATOM	6870	CD1	LEU	222	-16.513	-68.600	161.478	1.00 18.47	D	C
35	MOTA	6871	CD2	LEU	222	-17.900	-66.577	162.031	1.00 17.33	D	C
	MOTA	6872	С	LEU	222	-17.593	-65.454	157.771	1.00 21.62	D	C
	ATOM	6873	0	LEU	222	-17.955	-65.928	156.697	1.00 21.70	D	0
	MOTA	6874	N	ARG	223	-16.701	-64.480	157.871	1.00 21.04	D	N
	ATOM	6875	CA	ARG	223	-16.072	-63.870	156.709	1.00 21.51	D	С
40	MOTA	6876	СВ	ARG	223			156.424	1.00 22.86	D	C
	MOTA	6877	CG	ARG	223			155.282	1.00 25.93	D	C
	MOTA	6878	CD	ARG	223			155.770	1.00 27.48	D	С
	ATOM	6879	NE	ARG	223			156.520	1.00 28.36	D	N
	MOTA	6880	CZ	ARG	223			157.850	1.00 29.02	D	С
45	MOTA	6881	NH1	ARG	223			158.613	1.00 28.03	D	N
	MOTA	6882	NH2		223			158.425	1.00 28.61	D	N
	MOTA	6883	C	ARG	223			157.043	1.00 20.57	D	C
	MOTA	6884	0	ARG	223	-14.249	-62.850	157.889	1.00 20.75	D	0
	MOTA	6885	N	TYR	224	-13.722	-64.466	5 156.407	1.00 18.81	D	N
50	MOTA	6886		TYR	224			3 156.640	1.00 17.92	D	С
	MOTA	6887	CB	TYR	224			3 156.654		D	C
	MOTA	6888		TYR	224			3 157.744		D	C
	MOTA	6889		LTYR	224			2 157.530		D	С
	ATOM	6890	CE	LTYR	224			7 158.524		D	C
55	ATOM	6891		YYR	224			2 158.981		D	С
	MOTA	6892		2 TYR	224			7 159.985		D	С
	ATOM	6893		TYR	224			7 159.749		D	
	MOTA	6894	OH	TYR	224	-13.512	-69.25	6 160.725	1.00 15.77	D	0

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	MOTA	6895	С	TYR	224	-11.609	-63.534	155.557	1.00 17.55	D	С
	MOTA	6896	O	TYR	224		-63.721		1.00 16.13	D	0
	MOTA	6897	N	THR	225		-62.638		1.00 16.25	D	N
	MOTA	6898	CA	THR	225		-61.751		1.00 16.87	D	С
5	ATOM	6899	СВ	THR	225		-60.266		1.00 17.79	D	Ċ
	MOTA	6900	OG1	THR	225		-59.964		1.00 17.11	D	ō
	ATOM	6901	CG2	THR	225		-59.993		1.00 18.41	D	Č
	ATOM	6902	C	THR	225		-61.926		1.00 16.77	D	Ċ
	ATOM	6903	ō	THR	225		-62.593		1.00 15.99	D	Ō
10	ATOM	6904	N	ILE	226		-61.287		1.00 16.23	D	N
	ATOM	6905	CA	ILE	226		-61.354		1.00 16.57	D	C
	ATOM	6906	СВ	ILE	226		-60.731		1.00 17.02	D	č
	ATOM	6907		ILE	226		-59.202		1.00 15.14	D	C
	MOTA	6908	CG1		226		-61.149		1.00 16.61	D	Č
15	ATOM	6909	CD1		226		-60.691		1.00 14.79	D	Ċ
	MOTA	6910	C	ILE	226		-60.642		1.00 17.24	D	Č
	ATOM	6911	Ō	ILE	226		-60.956		1.00 16.91	D	Ō
	ATOM	6912	N	GLU	227		-59.705		1.00 16.07	D	N
	ATOM	6913	CA	GLU	227		-59.006		1.00 16.95	D	C
20	ATOM	6914	СВ	GLU	227		-57.820		1.00 16.63	D	Ċ
	ATOM	6915	CG	GLU	227			156.571	1.00 16.07	D	Č
	ATOM	6916	CD	GLU	227			155.195	1.00 16.66	D	č
	ATOM	6917	OE1		227				1.00 16.07	D	o
	ATOM	6918		GLU	227			154.187	1.00 17.97	D	ō
25	MOTA	6919	C	GLU	227			158.280	1.00 16.47	D	Ċ
	MOTA	6920	0	GLU	227			159.159	1.00 16.40	D	O
	MOTA	6921	N	ASP	228			158.309	1.00 15.66	D	N
	MOTA	6922	CA	ASP	228			159.393	1.00 16.51	D	C
	MOTA	6923	CB	ASP	228			159.311	1.00 15.72	D	Č
30	ATOM	6924	CG	ASP	228			159.555	1.00 16.63	D	Ċ
	MOTA	6925		ASP	228			160.540	1.00 16.26	D	ō
	ATOM	6926		ASP	228			158.768	1.00 16.57	D	ō
	ATOM	6927	С	ASP	228			159.361	1.00 15.84	D	С
	ATOM	6928	0	ASP	228	-4.791	-62.970	160.406	1.00 17.26	D	Ō
35	ATOM	6929	N	GLY	229			158.164	1.00 15.54	D	N
_	ATOM	6930	CA	GLY	229	-3.629	-63.735	158.053	1.00 14.97	D	C
	ATOM	6931	С	GLY	229			158.385	1.00 15.19	D	С
	MOTA	6932	0	GLY	229			159.010	1.00 14.72	D	0
	ATOM	6933	N	ALA	230			157.973	1.00 13.93	D	N
40	ATOM	6934	CA	ALA	230			158.233	1.00 14.50	D	С
	MOTA	6935	СВ	ALA	230	-1.844	-59.279	157.456	1.00 14.94	D	С
	MOTA	6936	С	ALA	230	-1.445	-60.268	159.732	1.00 14.36	D	С
	ATOM	6937	0	ALA	230	-0.341	-60.150	160.263	1.00 12.75	D	0
	MOTA	6938	N	ARG	231	-2.588	-60.164	160.410	1.00 14.18	D	N
45	ATOM	6939	CA	ARG	231	-2.596	-59.877	161.840	1.00 15.32	D	С
	ATOM	6940	CB	ARG	231	-4.014	-59.504	162.324	1.00 15.52	D	С
	MOTA	6941	CG	ARG	231	-4.594	-58.198	161.761	1.00 18.02	D	С
	MOTA	6942	CD	ARG	231	-3.828	-56.948	162.224	1.00 19.48	D	С
	MOTA	6943	NE	ARG	231	-3.939	-56.687	163.665	1.00 21.35	D	N
50	MOTA	6944	CZ	ARG	231	-4.921	-55.994	164.246	1.00 22.41	D	С
	ATOM	6945	NH1	ARG	231			163.527	1.00 20.52	D	N
	MOTA	6946	NH2	. ARG	231	-4.911	55.809	165.559	1.00 23.57	D	N
	ATOM	6947	С	ARG	231	-2.044	-61.026	162.698	1.00 14.73	D	C
	MOTA	6948	0	ARG	231	-1.608	-60.781	163.805	1.00 14.28	D	0
55	MOTA	6949	N	VAL	232			162.220	1.00 14.81	D	N
	ATOM	6950	CA	VAL	232			163.038	1.00 15.74	D	C
	MOTA	6951	СВ	VAL	232	-2.273	-64.700	162.869	1.00 16.44	D	С
	MOTA	6952	CG1	L VAL	232	-3.736	-64.514	163.233	1.00 17.47	D	

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	MOTA	6953	CG2		232			161.447	1.00 16.56	D	C
	ATOM	6954	С	VAL	232			162.744	1.00 15.66	D	С
	ATOM	6955	0	VAL	232			163.302	1.00 14.93	D	0
_	ATOM	6956	N	GLY	233	-		161.868	1.00 15.13	D	N
5	MOTA	6957	CA	GLY	233			161.586	1.00 15.93	D	С
	MOTA	6958	С	GLY	233			160.218	1.00 16.55	D	С
	MOTA	6959	0	GLY	233			159.961	1.00 16.84	D	0
	MOTA	6960	N	PHE	234	1.555	-63.741	159.347	1.00 16.39	D	N
	ATOM	6961	CA	PHE	234	2.008	-64.144	158.017	1.00 17.11	D	С
10	MOTA	6962	CB	PHE	234	0.882	-64.815	157.226	1.00 16.36	D	С
	MOTA	6963	CG	PHE	234	0.537	-66.194	157.703	1.00 16.05	D	С
	MOTA	6964	CD1	PHE	234			158.386	1.00 14.44	D	С
	MOTA	6965	CD2	PHE	234	1.382	-67.273	157.420	1.00 15.78	D	С
	MOTA	6966	CE1	PHE	234	-0.998	-67.724	158.779	1.00 16.61	D	С
15	MOTA	6967	CE2	PHE	234	1.043	-68.570	157.807	1.00 15.23	D	С
	MOTA	6968	CZ	PHE	234	-0.150	-68.799	158.486	1.00 15.89	D	С
	MOTA	6969	С	PHE	234	2.485	-62.930	157.234	1.00 18.16	D	С
	MOTA	6970	0	PHE	234	1.902	-61.844	157.340	1.00 18.39	D	0
	ATOM	6971	N	GLN	235	3.547	-63.112	156.451	1.00 19.12	D	N
20	ATOM	6972	CA	GLN	235	4.080	-62.032	155.624	1.00 19.59	D	С
	ATOM	6973	CB	GLN	235	5.445	-62.396	155.052	1.00 21.47	D	С
	ATOM	6974	CG	GLN	235	6.546	-62.520	156.080	1.00 25.04	D	С
	ATOM	6975	CD	GLN	235			155.433	1.00 27.60	D	С
	ATOM	6976		GLN	235			154.643	1.00 28.65	D	0
25	ATOM	6977	NE2		235			155.760	1.00 28.31	D	N
	ATOM	6978	C	GLN	235			154.476	1.00 19.54	D	C
	ATOM	6979	Ö	GLN	235			153.920	1.00 18.79	D	ō
	ATOM	6980	N	VAL	236			154.119	1.00 19.32	D	N
	ATOM	6981	CA	VAL	236			153.053	1.00 19.50	D	c
30	ATOM	6982	CB	VAL	236			152.866	1.00 19.33	D	c
30	ATOM	6983		VAL	236			151.660	1.00 13.53	D	Ċ
	ATOM	6984		VAL	236			154.126	1.00 18.96	D	C
	ATOM	6985	C	VAL	236			151.708	1.00 19.71	D	C
	ATOM	6986	Ö	VAL	236			151.708	1.00 19.46	D	ŏ
35	ATOM	6987	N	GLU	237			151.030	1.00 19.40	D	N
33	ATOM	6988	CA	GLU	237			150.014	1.00 20.19	D	C
	ATOM							149.802	1.00 21.32		C
		6989	CB	GLU	237					D	
	MOTA MOTA	6990	CG	GLU	237			148.485	1.00 27.69 1.00 31.29	D	C
40	_	6991	CD	GLU GLU	237					D	C
40	ATOM	6992			237 237			147.778 148.229	1.00 33.68 1.00 32.41	D D	0
	ATOM	6993		GLU							-
	ATOM			GLU	237			149.960			
	ATOM	6995	0	GLU	237			148.942	1.00 19.45	D	0
15	ATOM	6996	N	PHE	238			151.061	1.00 18.94	D	N
45	MOTA	6997	CA	PHE	238			151.152	1.00 18.95	D	C
	MOTA	6998	CB	PHE	238			152.511	1.00 18.87	D	C
	ATOM	6999	CG	PHE	238			2 152.862	1.00 18.24	D	C
	ATOM	7000		PHE	238			152.239	1.00 18.39	D	C
	MOTA	7001		PHE	238			153.797	1.00 18.85	D	C
50	MOTA	7002		PHE	238			152.545	1.00 18.05	D	C
	MOTA	7003	CE2		238			154.109	1.00 19.77	D	C
	MOTA	7004	CZ	PHE	238			153.480	1.00 18.89	D	С
	MOTA	7005	С	PHE	238			150.998	1.00 19.39	D	C
	MOTA	7006	0	PHE	238			1 150.239	1.00 18.49	D	
55	MOTA	7007	N	LEU	239			3 151.713	1.00 18.74	D	
	MOTA	7008	CA	LEU	239			5 151.675	1.00 19.82	D	_
	MOTA	7009	CB	LEU	239			2 152.611	1.00 18.63	D	_
	ATOM	7010	CG	LEU	239	-1.538	-63.153	3 153.992	1.00 19.49	D	C

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	ATOM	7011	CD1		239		-64.510		1.00 16.15	D	C
	MOTA	7012	CD2		239		-62.049		1.00 16.47	D	C
	MOTA	7013		LEU	239		-63.697		1.00 20.21	D	C
E	MOTA	7014		LEU	239		-64.315		1.00 19.21	D	0
5	MOTA	7015		GLU	240		-62.755		1.00 20.94	D	N
	ATOM	7016		GLU	240		-62.413		1.00 22.99	D	C
	MOTA	7017		GLU	240		-61.255		1.00 25.70	D	С
	MOTA	7018		GLU	240		-59.897		1.00 28.57	D	C
40	MOTA	7019		GLU	240		-59.334		1.00 32.06	D	C
10	MOTA	7020		GLU	240		-60.068		1.00 33.41	D	0
	MOTA	7021		GLU	240		-58.151		1.00 34.50	D	0
	MOTA	7022	C	GLU	240		-63.633		1.00 22.86	D	C
	ATOM	7023	0	GLU	240		-63.874		1.00 22.97	D	0
15	ATOM	7024	N	LEU	241		-64.400		1.00 22.57	D	N
15	ATOM	7025	CA	LEU	241		-65.603		1.00 23.67	D	C
	ATOM	7026	CB	LEU	241		-66.270		1.00 25.60	D	C
	MOTA	7027	CG	LEU	241		-67.315		1.00 28.83	D	C
	ATOM	7028	CD1		241		-68.588		1.00 30.65	D	C
20	MOTA	7029	CD2		241		-66.775 -66.551		1.00 29.93	D	C
20	MOTA	7030 7031	C	LEU	241				1.00 22.49	D	C
	ATOM	7031	O N	LEU	241 242		-67.113 -66.692		1.00 21.13	D	0
	ATOM ATOM	7032	CA	LEU	242			148.201	1.00 21.14 1.00 20.53	D	N
	ATOM	7033	CB	LEU	242			150.114	1.00 20.33	D	C
25	ATOM	7035	CG	LEU	242			150.114	1.00 21.39	D	C
20	ATOM	7036		LEU	242			150.701	1.00 22.85	D	C
	ATOM	7030		LEU	242			152.211	1.00 22.86	D D	C
	ATOM	7037	C	LEU	242			148.017	1.00 22.33	מ	c
	ATOM	7039	Ö	LEU	242			147.438	1.00 20.13	D	Ö
30	ATOM	7040	N	PHE	243			148.160	1.00 19.21	D	N
00	MOTA	7041	CA	PHE	243			147.646	1.00 19.10	D	C
	ATOM	7042	CB	PHE	243			148.242	1.00 17.31	D	Č
	ATOM	7043	CG	PHE	243			149.650	1.00 16.71	D	Ċ
	ATOM	7044		PHE	243			150.713	1.00 15.59	D	Č
35	ATOM	7045	CD2		243			149.905	1.00 16.40	D	č
	ATOM	7046		PHE	243			152.009	1.00 15.90	D	č
	ATOM	7047		PHE	243			151.204	1.00 16.02	D	Ċ
	ATOM	7048	CZ	PHE	243			152.254	1.00 15.02	D	Ċ
	MOTA	7049	C	PHE	243			146.117	1.00 19.44	D	Č
40	MOTA	7050	0	PHE	243			145.533	1.00 18.99	D	Ō
	MOTA	7051	N	HIS	244			145.474	1.00 19.69	D	N
	ATOM	7052	CA	HIS	244	-3.804	-65.152	144.016	1.00 20.80	D	С
	MOTA	7053	CB	HIS	244	-2.404	-64.783	143.505	1.00 22.74	D	C
	ATOM	7054	CG	HIS	244	-2.235	-64.959	142.025	1.00 26.38	D	С
45	ATOM	7055	CD2	HIS	244	-2.396	-64.088	140.999	1.00 27.16	D	С
	MOTA	7056	ND1	HIS	244	-1.905	-66.170	141.448	1.00 27.90	D	N
	MOTA	7057		HIS	244	-1.873	-66.036	140.134	1.00 27.13	D	C
	ATOM	7058	NE2	HIS	244	-2.167	-64.783	139.836	1.00 27.37	D	N
	MOTA	7059	C	HIS	244			143.591	1.00 19.50	D	C
50	MOTA	7060	0	HIS	244			142.621	1.00 18.50	D	0
	ATOM	7061	N	PHE	245			144.326	1.00 18.62	D	N
	MOTA	7062	CA	PHE	245			144.036	1.00 17.80		C
	MOTA	7063	СВ	PHE	245			145.055	1.00 17.14		C
	MOTA	7064	CG	PHE	245			145.069	1.00 16.75		C
55	MOTA	7065		PHE	245			144.105	1.00 16.57		С
	ATOM	7066		PHE	245			146.007	1.00 16.20		C
	MOTA	7067		PHE	245			144.068	1.00 16.23		C
	MOTA	7068	CE2	PHE	245	-5.372	-72.884	145.984	1.00 16.41	D	С

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	MOTA	7069	CZ	PHE	245	-4.992	-73.812	145.009	1.00 15.98	D	С
	MOTA	7070	С	PHE	245	-5.430	-69.170	144.100	1.00 17.44	D	С
	MOTA	7071	0	PHE	245	-6.005	-69.787	143.212	1.00 17.41	D	0
	MOTA	7072	N	HIS	246	-6.068	-68.669	145.154	1.00 16.27	D	N
5	ATOM	7073	CA	HIS	246	-7.505	-68.853	145.316	1.00 16.76	D	С
	ATOM	7074	СВ	HIS	246	-7.939	-68.409		1.00 14.87	D	С
	ATOM	7075	CG	HIS	246		-69.466		1.00 15.84	D	C
	ATOM	7076	CD2		246		-69.620		1.00 14.94	D	Ċ
	ATOM	7077	ND1		246		-70.574		1.00 13.95	D	N
10	MOTA	7078	CE1		246		-71.366		1.00 14.24	D	C
	ATOM	7079		HIS	246		-70.810		1.00 14.24	D	Ŋ
	ATOM	7080	C	HIS	246		-68.158				C
	ATOM	7081	0		246				1.00 16.72	D	
				HIS				143.694	1.00 16.62	D	0
4.5	ATOM	7082	N	GLY	247			143.940	1.00 16.82	D	N
15	ATOM	7083	CA	GLY	247			142.907	1.00 18.06	D	С
	MOTA	7084	C	GLY	247			141.569	1.00 18.01	D	С
	MOTA	7085	0	GLY	247			140.867	1.00 18.60	D	0
	MOTA	7086	N	THR	248			141.222	1.00 18.13	D	N
	MOTA	7087	CA	THR	248			139.953	1.00 18.70	D	С
20	MOTA	7088	CB	THR	248				1.00 18.46	D	С
	MOTA	7089	OG1	THR	248	-4.933	-67.084	139.926	1.00 19.05	D	0
	MOTA	7090	CG2	THR	248	-5.389	-68.857	138.368	1.00 18.69	D	С
	MOTA	7091	C	THR	248	-7.937	-69.321	139.878	1.00 18.91	D	С
	MOTA	7092	0	THR	248	-8.565	-69.621	138.872	1.00 18.39	D	0
25	MOTA	7093	N	LEU	249	-7.882	-70.110	140.943	1.00 19.20	D	N
	ATOM	7094	CA	LEU	249	-8.593	-71.378	140.969	1.00 20.23	D	С
	ATOM	7095	СВ	LEU	249			142.259	1.00 19.48	D	C
	MOTA	7096	CG	LEU	249		-73.523		1.00 20.29	D	C
	MOTA	7097		LEU	249			141.232	1.00 20.40	D	Č
30	ATOM	7098		LEU	249			143.713	1.00 18.44	Ď	č
00	MOTA	7099	C	LEU	249			140.867	1.00 20.65	D	C
	MOTA	7100	Ö	LEU	249			140.109	1.00 20.03	D	0
	ATOM	7101	N	ARG	250			141.621	1.00 13.05	D	N
	ATOM	7101	CA	ARG	250			141.626	1.00 21.13	D	C
35		7102	CB	ARG	250			142.662	1.00 24.41		
33	MOTA	7103	CG	ARG	250			142.8835	1.00 24.41	D	C
	ATOM							143.633		D	C
	MOTA	7105	CD	ARG	250				1.00 27.83	D	С
	ATOM	7106	NE	ARG	250			144.495	1.00 29.89	D	N
40	MOTA	7107	CZ	ARG	250			144.354	1.00 32.93	D	С
40	ATOM	7108		ARG	250			145.412	1.00 35.07	D	N
	ATOM	7109		ARG	250			143.159	1.00 33.97	D	N
	MOTA	7110	C	ARG	250			140.284	1.00 23.27	D	C
	MOTA	7111	0	ARG	250			139.972	1.00 21.85	D	0
	MOTA	7112	N	LYS	251			139.504	1.00 24.10	D	N
45	MOTA	7113	CA	LYS	251			138.211	1.00 25.70	D	C
	MOTA	7114	CB	LYS	251			137.640	1.00 26.54	D	С
	ATOM	7115	CG	LYS	251			138.421	1.00 27.85	D	С
	MOTA	7116	CD	LYS	251	-10.374	-64.842	137.821	1.00 30.19	D	C
	ATOM	7117	CE	LYS	251			138.716	1.00 31.63	D	C
50	ATOM	7118	NZ	LYS	251	-9.159	-62.675	138.208	1.00 34.50	D	N
	MOTA	7119	С	LYS	251			137.190	1.00 26.18	D	С
	MOTA	7120	Ō	LYS	251			136.192	1.00 26.51	D	ō
	ATOM	7121	N	LEU	252			137.431	1.00 26.18	D	N
	ATOM	7122	CA	LEU	252			3 136.509	1.00 26.53	D	C
55	ATOM	7123	СВ	LEU	252			136.745	1.00 24.32	D	Ċ
-	ATOM	7124	CG	LEU	252			136.495	1.00 23.79	D	C
	ATOM	7124		LEU	252			136.493	1.00 21.82	D	C
	ATOM	7125		FEO FEO	252			135.786	1.00 21.82	D	C
	ATOM	1120	CDA	טפע י	2.14	-9.3/0	-/1.001	133.034	1.00 22.08	ע	C

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	MOTA	7127	С	LEU	252	-13.446	-72.259	136.628	1.00 27.56	D	С
	MOTA	7128	0	LEU	252	-13.794	-73.108	135.812	1.00 28.31	D	0
	ATOM	7129	N	GLN	253	-14.222	-71.868	137.636	1.00 28.57	D	N
	MOTA	7130	CA	GLN	253	-15.556	-72.421	137.863	1.00 29.80	D	С
5	MOTA	7131	CB	GLN	253	-16.554	-71.824	136.862	1.00 31.77	D	С
	ATOM	7132	CG	GLN	253	-16.555	-70.298	136.817	1.00 35.28	D	С
	ATOM	7133	CD	GLN	253	-17.738	-69.711	136.048	1.00 38.18	D	С
	MOTA	7134	OE1		253		-68.597		1.00 39.03	D	0
	ATOM	7135	NE2		253	-18.847	-70.451	135.992	1.00 39.86	D	N
10	ATOM	7136	С	GLN	253	-15.576	-73.942	137.752	1.00 29.42	D	C
	MOTA	7137	0	GLN	253		-74.502		1.00 29.12	D	Ō
	ATOM	7138	N	LEU	254		-74.609		1.00 28.93	D	N
	ATOM	7139	CA	LEU	254		-76.065		1.00 28.81	D	C
	ATOM	7140	СВ	LEU	254		-76.572		1.00 26.92	D	C
15	ATOM	7141	CG	LEU	254		-76.144		1.00 26.12	Ď	C
. •	ATOM	7142		LEU	254		-76.798		1.00 23.47	D	C
	ATOM	7143		LEU	254		-76.530		1.00 25.72	D	C
	ATOM	7144	C	LEU	254		-76.687		1.00 29.64	D	C
	ATOM	7145	Ö	LEU	254		-76.068		1.00 29.37	D	0
20	ATOM	7146	N	GLN	255		-77.926		1.00 29.37	D	И
20	ATOM	7147	CA	GLN	255		-78.681		1.00 30.18		C
	ATOM	7148	CB	GLN	255		-79.489			D	C
	ATOM	7149	CG	GLN	255		-79.469 -78.843		1.00 34.30 1.00 38.01	D	
	ATOM	7150	CD	GLN	255		-79.210		1.00 38.01	D	C
25										D	C
23	MOTA	7151	OE1		255		-80.360		1.00 42.15	D	0
	MOTA	7152	NE2		255		-78.230		1.00 42.18	D	N
	ATOM	7153	C	GLN	255			140.251	1.00 31.09	D	С
	MOTA	7154	0	GLN	255			140.292	1.00 29.84	D	0
20	ATOM	7155	N	GLU	256			141.127	1.00 30.54	D	N
30	MOTA	7156	CA	GLU	256			142.219	1.00 30.46	D	C
	MOTA	7157	CB	GLU	256			142.920	1.00 32.16	D	C
	ATOM	7158	CG	GLU	256			143.620	1.00 34.89	D	С
	MOTA	7159	CD	GLU	256			144.663	1.00 35.98	D	С
0.5	MOTA	7160	OE1		256			144.320	1.00 36.28	D	0
35	MOTA	7161		GLU	256			145.820	1.00 36.73	D	0
	MOTA	7162	С	GLU	256			141.861	1.00 29.44	D	С
	MOTA	7163	0	GLU	256			142.486	1.00 29.52	D	0
	MOTA	7164	N	PRO	257			140.862	1.00 28.35	D	N
	MOTA	7165	CD	PRO	257			140.045	1.00 29.24	D	С
40	MOTA	7166	CA	PRO	257	-15.605	-83.927	140.526	1.00 27.52	D	С
	MOTA	7167	CB	PRO	257			139.347	1.00 27.78	D	C
	MOTA	7168	CG	PRO	257	-17.266	-83.672	138.815	1.00 29.42	D	С
	MOTA	7169	С	PRO	257	-14.197	-83.416	140.215	1.00 25.67	D	С
	MOTA	7170	0	PRO	257	-13.212	-84.074	140.539	1.00 24.99	D	0
45	MOTA	7171	N	GLU	258	-14.098	-82.247	139.590	1.00 24.23	D	N
	ATOM	7172	CA	GLU	258	-12.795	-81.671	139.273	1.00 22.93	D	С
	MOTA	7173	CB	GLU	258	-12.969	-80.476	138.329	1.00 21.97	D	C
	ATOM	7174	CG	GLU	258			136.971	1.00 23.09	D	C
	MOTA	7175	CD	GLU	258			136.109	1.00 22.64	D	C
50	MOTA	7176	OE1	GLU	258			136.653	1.00 22.76	D	O
	MOTA	7177	OE2		258			134.876	1.00 22.39	D	ō
	ATOM	7178	C	GLU	258			140.568	1.00 22.29	D	Č
	MOTA	7179	ō	GLU	258			140.733	1.00 21.58	D	ō
	ATOM	7180	N	TYR	259			141.484	1.00 20.81	D	N
55	ATOM	7181	CA	TYR	259			142.762	1.00 20.66	D	C
	ATOM	7182	СВ	TYR	259			143.635	1.00 20.00	D	C
	MOTA	7183	CG	TYR	259			143.490	1.00 19.41	D	C
	ATOM	7184		L TYR	259			143.490	1.00 18.07	D	
	AIUM	1104	CD	LIIK	439	-12.1/0	-//.210	143.023	1.00 18.03	ע	С

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	MOTA	7185	CE1	TYR	259	-12.205			1.00 16.82	D	С
	MOTA	7186	CD2	TYR	259	-14.429			1.00 17.04	D	С
	MOTA	7187	CE2	TYR	259	-14.478	-75.954	142.940	1.00 16.86	D	С
_	MOTA	7188	CZ	TYR	259	-13.363			1.00 17.78	D	С
5	MOTA	7189	OH	TYR	259	-13.426			1.00 16.55	D	0
	ATOM	7190	С	TYR	259	-11.740			1.00 20.57	D	С
	MOTA	7191	0	TYR	259	-10.625			1.00 19.89	D	0
	MOTA	7192	N	VAL	260	-12.568			1.00 20.84	D	N
	MOTA	7193	CA	VAL	260	-12.169			1.00 21.82	D	С
10	MOTA	7194	CB	VAL	260	-13.386			1.00 23.38	D	С
	MOTA	7195		VAL	260	-13.761			1.00 23.89	D	С
	MOTA	7196	CG2	VAL	260			145.719	1.00 24.50	D	С
	MOTA	7197	С	VAL	260			143.679	1.00 21.20	D	С
	MOTA	7198	0	VAL	260			144.373	1.00 21.64	D	0
15	MOTA	7199	N	LEU	261			142.353	1.00 20.69	D	N
	MOTA	7200	CA	LEU	261			141.656	1.00 21.07	D	С
	MOTA	7201	СВ	LEU	261			140.158	1.00 21.76	D	С
	MOTA	7202	CG	LEU	261			139.803	1.00 21.71	D	С
	ATOM	7203		LEU	261			138.346	1.00 21.95	D	С
20	MOTA	7204		LEU	261			140.090	1.00 20.53	D	С
	MOTA	7205	С	LEU	261			141.897	1.00 21.22	D	С
	MOTA	7206	0	LEU	261			141.996	1.00 21.66	D	0
	MOTA	7207	N	LEU	262			141.999	1.00 21.22	D	N
	MOTA	7208	CA	LEU	262			142.266	1.00 21.43	D	С
25	MOTA	7209	СВ	LEU	262			142.212	1.00 23.16	D	С
	ATOM	7210	CG	LEU	262			141.743	1.00 25.15	D	С
	MOTA	7211		LEU	262			140.390	1.00 24.87	D	C
	MOTA	7212		LEU	262			141.647	1.00 26.33	D	С
	MOTA	7213	С	LEU	262			143.663	1.00 21.12	D	С
30	MOTA	7214	0	LEU	262			143.877	1.00 20.89	D	0
	MOTA	7215	N	ALA	263			144.608	1.00 19.39	D	N
	MOTA	7216	CA	ALA	263			145.972	1.00 19.66	D	C
	MOTA	7217	CB	ALA	263			146.865	1.00 18.64	D	C
0.5	MOTA	7218	C	ALA	263			145.989	1.00 19.63	D	C
35	MOTA	7219	0	ALA	263			146.686	1.00 19.49	D	0
	MOTA	7220	N	ALA	264			145.216	1.00 19.23	D	N
	MOTA	7221	CA	ALA	264			145.136	1.00 19.14	D	C
	ATOM	7222	СВ	ALA	264			144.300	1.00 18.99	D	C
40	ATOM	7223	C	ALA	264			144.518	1.00 19.43	D	C
40	ATOM	7224	0	ALA	264			. 144.940	1.00 18.52	D	0
	ATOM	7225	N	MET MET	265 265			7 143.517 2 142.881	1.00 18.79 1.00 20.11	D	N C
	ATOM	7226								D	_
	ATOM	7227	CB	MET	265			5 141.642 2 140.467	1.00 21.94 1.00 23.98	D	C
45	ATOM	7228	CG	MET	265					D	C
45	MOTA	7229	SD CE	MET	265 265			2 139.013 1 138.431	1.00 27.16 1.00 25.04	D	S
	MOTA	7230 7231		MET				7 143.868		D	
	ATOM		C	MET	265				1.00 19.34 1.00 18.51	D	C
	MOTA	7232	0	MET	265 266			5 143.850 2 144.732	1.00 18.31	D	0
50	MOTA	7233	N	ALA	266			5 145.745	1.00 19.42	D	
50	ATOM ATOM	7234 7235	CA CB	ALA ALA	266			9 145.745		D D	
		7236	СВ	ALA	266			2 146.762		D	
	MOTA				266			9 147.200			
	ATOM	7237		ALA LEU	267			147.200 1 147.125			
55	ATOM	7238 7239		LEU	267			3 148.080			
55	ATOM			LEU	267			5 148.080 5 148.225			
	ATOM	7240		LEU	267			9 149.565			-
	MOTA	7241		LEU LEU	267			4 149.295			
	MOTA	7242	CD.	rneu	201	-5.455	-07.23	H 147.675	1.00 21.48	ט	C

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		7043	000		0.68	2 560	00 000	150 440	1 00 00 05	_	_
	MOTA	7243	CD2		267		-88.082		1.00 22.95	D	C
	MOTA	7244	С	LEU	267		-87.564	-	1.00 23.83	D	C
	MOTA	7245	0	LEU	267		-87.902		1.00 23.92	D	0
_	MOTA	7246	N	PHE	268		-88.050		1.00 25.59	D	N
5	MOTA	7247	CA	PHE	268		-89.066		1.00 27.95	D	С
	MOTA	7248	CB	PHE	268		-90.039		1.00 26.41	D	С
	MOTA	7249	CG	PHE	268	-3.071	-90.769	145.657	1.00 25.14	D	С
	ATOM	7250	CD1	PHE	268	-4.387	-90.525	145.286	1.00 24.11	D	С
	ATOM	7251	CD2	PHE	268	-2.822	-91.688	146.676	1.00 25.38	D	С
10	ATOM	7252	CE1	PHE	268	-5.438	-91.176	145.916	1.00 23.69	D	С
	ATOM	7253	CE2		268		-92.351		1.00 23.66	D	C
	ATOM	7254	CZ	PHE	268		-92.093		1.00 23.71	D	C
	ATOM	7255	C	PHE	268			145.222	1.00 30.43	D	Ċ
	ATOM	7256	ō	PHE	268			144.057	1.00 30.67	D	ŏ
15	ATOM	7257	N	SER	269			145.948	1.00 33.96	D	N
10	MOTA	7258	CA	SER	269			145.422	1.00 37.03		C
	ATOM	7259	CB	SER	269			145.422		D	C
									1.00 37.39	D	
	MOTA	7260	OG	SER	269			145.463	1.00 38.30	D	0
00	MOTA	7261	С	SER	269			145.896	1.00 39.10	D	C
20	MOTA	7262	0	SER	269			147.096	1.00 38.72	D	0
	ATOM	7263	N	PRO	270			144.954	1.00 41.49	D	Ŋ
	ATOM	7264	CD	PRO	270			143.499	1.00 41.60	D	C
	ATOM	7265	CA	PRO	270	4.884	-89.957	145.270	1.00 43.13	D	C
	MOTA	7266	CB	PRO	270	5.051	-90.704	143.951	1.00 42.69	D	C
25	MOTA	7267	CG	PRO	270	4.821	-89.621	142.940	1.00 42.36	D	С
	ATOM	7268	С	PRO	270	6.211	-89.397	145.782	1.00 44.56	D	С
	ATOM	7269	0	PRO	270	7.015	-90.131	146.361	1.00 45.15	D	0
	ATOM	7270	N	ASP	271			145.577	1.00 45.55	D	N
	MOTA	7271	CA	ASP	271			146.010	1.00 46.83	D	C
30	ATOM	7272	СВ	ASP	271			144.996	1.00 48.20	D	c
00	ATOM	7273	CG	ASP	271			144.852	1.00 49.90	D	Ċ
	ATOM	7274		ASP	271			144.759	1.00 50.47	D	o
	ATOM	7275		ASP	271			144.818	1.00 50.92		
					271					D	0
35	ATOM	7276	C	ASP				147.418	1.00 47.00	D	C
33	ATOM	7277	0	ASP	271			147.815	1.00 47.50	D	0
	MOTA	7278	N	ARG	272			148.172	1.00 46.37	D	Ŋ
	MOTA	7279	CA	ARG	272			149.534	1.00 45.93	D	C
	MOTA	7280	CB	ARG	272			150.050	1.00 45.63	D	С
4.0	ATOM	7281	CG	ARG	272			150.556	1.00 44.83	D	С
40	ATOM	7282	CD	ARG	272			149.482	1.00 42.90	D	С
	ATOM	7283	NE	ARG	272			149.425	1.00 41.67	D	N
	ATOM	7284	CZ	ARG	272			148.936	1.00 40.40	D	С
	MOTA	7285		ARG	272	1.366	-83.563	148.445	1.00 38.84	D	N
	ATOM	7286	NH2	ARG	272	2.770	-81.826	148.939	1.00 40.16	D	N
45	MOTA	7287	С	ARG	272	7.501	-87.396	150.435	1.00 46.08	D	С
	ATOM	7288	0	ARG	272	7.880	-88.555	150.246	1.00 45.78	D	0
	ATOM	7289	N	PRO	273			151.428	1.00 46.22	D	N
	ATOM	7290	CD	PRO	273			151.712	1.00 45.76	D	C
	ATOM	7291	CA	PRO	273			152.334	1.00 46.41	D	C
50	ATOM	7292	СВ	PRO	273			153.132	1.00 46.21	D	c
00	ATOM	7293	CG	PRO	273			153.132	1.00 46.57		
		7294	C	PRO	273			153.133		D	C
	MOTA								1.00 46.69	D	C
	MOTA	7295	0	PRO	273			153.938	1.00 46.36	D	0
EE	MOTA	7296	N	GLY	274			153.193	1.00 46.90	D	N
55	MOTA	7297	CA	GLY	274			154.010	1.00 47.62	D	C
	MOTA	7298	C	GLY	274			153.314	1.00 48.15	D	С
	MOTA	7299	0	GLY	274			153.951	1.00 48.17	D	0
	MOTA	7300	N	VAL	275	7.748	-91.462	152.012	1.00 48.62	D	N

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	MOTA	7301		VAL	275		2
	MOTA	7302	-	VAL	275		C
	MOTA	7303	CG1		275		C
_	MOTA	7304	CG2		275		С
5	MOTA	7305	C	VAL	275		С
	MOTA	7306	0	VAL	275		0
	ATOM	7307	N	THR	276	7.136 -94.818 151.224 1.00 50.30 D I	N
	ATOM	7308	CA	THR	276	7.748 -96.112 150.957 1.00 51.08 D	C
	MOTA	7309	CB	THR	276	7.258 -97.168 151.975 1.00 51.15 D	С
10	MOTA	7310	OG1	THR	276	7.755 -96.838 153.277 1.00 51.62 D	0
	ATOM	7311	CG2	THR	276	7.742 -98.559 151.590 1.00 51.91 D	С
	MOTA	7312	С	THR	276	7.425 -96.590 149.542 1.00 51.59 D	С
	ATOM	7313	0	THR	276	8.321 -96.729 148.710 1.00 52.05 D	0
	ATOM	7314	N	GLN	277		N
15	MOTA	7315	CA	GLN	277		C
. •	ATOM	7316	СВ	GLN	277		c
	ATOM	7317	CG	GLN	277		c
	ATOM	7318	CD	GLN	277		c
	ATOM	7319	OE1		277		ō
20	MOTA	7320	NE2	GLN	277		N
20	ATOM	7321	C	GLN	277		C
	ATOM	7322	Ö	GLN	277		0
	ATOM	7323	N	ARG	278		N
	ATOM	7323	CA	ARG	278		C
25	ATOM	7324	CB	ARG	278		C
23	ATOM	7325	CG	ARG	278		C
		7327	CD	ARG	278		C
	MOTA		NE	ARG			
	MOTA	7328	CZ		278		N
30	MOTA	7329		ARG	278		C
30	ATOM	7330		ARG	278		N
	ATOM	7331		ARG	278		N
	MOTA	7332	C	ARG	278		C
	ATOM	7333	0	ARG	278		0
25	MOTA	7334	N	ASP	279		N
35	MOTA	7335	CA	ASP	279	5.982 -96.516 142.532 1.00 49.17 D	C
	ATOM	7336	CB	ASP	279	6.693 -97.818 142.151 1.00 51.29 D	С
	MOTA	7337	CG	ASP	279	8.109 -97.585 141.643 1.00 53.01 D	C
	MOTA	7338		ASP	279	8.825 -96.739 142.221 1.00 54.57 D	0
40	MOTA	7339		ASP	279	8.510 -98.254 140.667 1.00 54.28 D	0
40	MOTA	7340	С	ASP	279	4.470 -96.718 142.508 1.00 47.62 D	С
	MOTA	7341	0	ASP	279	3.801 -96.316 141.560 1.00 47.32 D	0
	MOTA	7342	N	GLU	280		N
	MOTA	7343	CA	GLU	280	2.495 -97.579 143.622 1.00 44.64 D	С
4-	MOTA	7344	CB	GLU	280	2.157 -98.466 144.826 1.00 45.89 D	С
45	MOTA	7345	CG	GLU	280	2.755 -99.866 144.754 1.00 48.45 D	C
	ATOM	7346	CD	GLU	280	4.074 -99.997 145.509 1.00 49.73 D	C
	MOTA	7347		GLU	280	4.986 -99.165 145.302 1.00 49.56 D	0
	ATOM	7348		GLU	280	4.199-100.947 146.312 1.00 51.25 D	0
	ATOM	7349	C	GLU	280	1.728 -96.260 143.715 1.00 42.74 D	С
50	MOTA	7350	0	GLU	280	0.715 -96.074 143.045 1.00 42.04 D	0
	MOTA	7351	N	ILE	281	2.213 -95.348 144.550 1.00 41.01 D	N
	MOTA	7352	CA	ILE	281	1.566 -94.053 144.706 1.00 40.02 D	С
	ATOM	7353	CB	ILE	281	2.166 -93.277 145.900 1.00 39.33 D	С
	MOTA	7354		ILE	281	1.636 -91.847 145.919 1.00 38.82 D	C
55	MOTA	7355	CG1	ILE	281	1.807 -93.997 147.206 1.00 38.81 D	С
	ATOM	7356	CD1	ILE	281	2.431 -93.392 148.443 1.00 37.90 D	С
	MOTA	7357	С	ILE	281	1.694 -93.231 143.420 1.00 39.83 D	С
	MOTA	7358	0	ILE	281	0.794 -92.466 143.069 1.00 39.50 D	0

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	MOTA	7359	N	ASP	282	2.810	-93.405	142.718	1.00 39.49	D	N
	MOTA	7360	CA	ASP	282	3.049	-92.698	141.465	1.00 39.92	D	С
	MOTA	7361	CB	ASP	282	4.440	-93.042	140.924	1.00 42.08	D	С
_	MOTA	7362	CG	ASP	282				1.00 44.33	D	С
5	MOTA	7363	OD1	ASP	282	4.385	-91.694	138.956	1.00 45.27	D	0
	MOTA	7364	OD2	ASP	282	6.053	-91.388	140.351	1.00 46.09	D	0
	MOTA	7365	С	ASP	282	1.977	-93.115	140.453	1.00 39.04	D	С
	ATOM	7366	0	ASP	282	1.403	-92.273	139.762	1.00 38.00	D	0
	MOTA	7367	N	GLN	283		-94.417		1.00 38.52	D	N
10	ATOM	7368	CA	GLN	283	0.691	-94.943	139.471	1.00 38.31	D	С
	MOTA	7369	СВ	GLN	283	0.725	-96.476	139.448	1.00 40.70	D	С
	MOTA	7370	CG	GLN	283		-	139.041	1.00 44.18	D	С
	MOTA	7371	CD	GLN	283			138.677	1.00 46.41	D	С
	MOTA	7372	OE1	GLN	283			139.404	1.00 47.67	D	0
15	MOTA	7373	NE2	GLN	283	2.546	-98.941	137.546	1.00 46.87	D	N
	ATOM	7374	С	GLN	283	-0.709	-94.475	139.860	1.00 36.85	D	С
	MOTA	7375	0	GLN	283	-1.567	-94.268	139.000	1.00 36.29	D	0
	MOTA	7376	N	LEU	284			141.159	1.00 35.69	D	N
	MOTA	7377	CA	LEU	284	-2.249	-93.865	141.624	1.00 34.25	D	С
20	MOTA	7378	CB	LEU	284	-2.346	-93.955	143.153	1.00 33.82	D	С
	MOTA	7379	CG	LEU	284	-2.472	-95.364	143.747	1.00 33.47	D	С
	MOTA	7380	CD1	LEU	284	-2.597	-95.281	145.259	1.00 32.83	D	С
	MOTA	7381	CD2	LEU	284	-3.696	-96.053	143.164	1.00 32.79	D	С
	MOTA	7382	С	LEU	284	-2.483	-92.430	141.171	1.00 33.26	D	С
25	MOTA	7383	0	LEU	284	-3.593	-92.069	140.781	1.00 31.51	D	0
	MOTA	7384	N	GLN	285	-1.435	-91.611	141.216	1.00 33.02	D	N
	MOTA	7385	CA	GLN	285	-1.570	-90.224	140.794	1.00 33.96	D	С
	ATOM	7386	CB	GLN	285	-0.281	-89.442	141.037	1.00 34.10	D	С
	ATOM	7387	CG	GLN	285	-0.430	-87.960	140.713	1.00 35.79	D	С
30	MOTA	7388	CD	GLN	285	0.792	-87.162	141.077	1.00 36.64	D	С
	MOTA	7389	OE1	GLN	285	1.848	-87.307	140.461	1.00 38.71	D	0
	MOTA	7390	NE2	GLN	285	0.665	-86.317	142.095	1.00 36.76	D	N
	MOTA	7391	С	GLN	285	-1.929	-90.141	139.317	1.00 33.52	D	C
	ATOM	7392	0	GLN	285	-2.799	-89.366	138.923	1.00 33.70	D	0
35	MOTA	7393	N	GLU	286	-1.258	-90.942	138.501	1.00 33.33	D	N
·	ATOM	7394	CA	GLU	286	-1.522	-90.936	137.069	1.00 34.29	D	C
	MOTA	7395	CB	GLU	286	-0.568	-91.902	136.358	1.00 36.73	D	C
	MOTA	7396	CG	GLU	286	0.880	-91.694	136.801	1.00 41.05	D	С
	ATOM	7397	CD	GLU	286	1.897	-92.388	135.919	1.00 44.06	D	C
40	ATOM	7398	OE1	GLU	286	1.725	-93.594	135.629	1.00 46.05	D	0
	ATOM	7399	OE2	GLU	286			135.525	1.00 45.70	D	0
	MOTA	7400	C	GLU	286	-2.976	-91.306	136.817	1.00 32.71	D	C
	ATOM	7401	0	GLU	286	-3.628	-90.726	135.951	1.00 31.80	D	0
	MOTA	7402	N	GLU	287			137.593	1.00 32.21	D	N
45	MOTA	7403	CA	GLU	287			137.467	1.00 31.48	D	C
	ATOM	7404	CB	GLU	287	-5.186	-93.841	138.421	1.00 33.15	D	С
	ATOM	7405	CG	GLU	287	-6.573	-94.430	138.208	1.00 36.26	D	С
	MOTA	7406	CD	GLU	287	-6.975	-95.435	139.278	1.00 38.20	D	С
	MOTA	7407	OE1	GLU	287	-6.145	-96.301	L 139.634	1.00 38.75	D	0
50	MOTA	7408	OE2	GLU	287	-8.131	95.363	3 139.752	1.00 38.76	D	0
	MOTA	7409	C	GLU	287	-5.796	-91.517	7 137.787	1.00 29.64	D	С
	MOTA	7410	0	GLU	287			3 137.122	1.00 30.00	D	0
	MOTA	7411	N	MET	288			3 138.818		D	N
	MOTA	7412	CA	MET	288	-6.258	-89.589	9 139.197			С
55	MOTA	7413	СВ	MET	288			5 140.464		D	C
	ATOM	7414	CG	MET	288	-5.579	-89.80	141.706	1.00 26.67	D	С
	MOTA	7415	SD	MET	288	-7.136	-90.50	1 142.292	1.00 26.45	D	S
	ATOM	7416	CE	MET	288	-6.849	92.26	4 141.971	1.00 27.72	D	С

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	ATOM	7417	С	MET	288	-6 207	-88.592	138 040	1.00 26.19	D	С
	ATOM	7418	ō	MET	288		-88.074		1.00 25.68	D	ō
	MOTA	7419	N	ALA	289		-88.348		1.00 25.78	D	N
	ATOM	7420	CA	ALA	289		-87.407		1.00 25.78	D	C
5	ATOM	7421	CB	ALA	289			136.108	1.00 24.39	D	C
9		7422	C	ALA	289		-87.746		1.00 25.97	D	C
	MOTA	7423	0	ALA	289		-86.883		1.00 25.73	D	0
	MOTA	7424		LEU	290		-89.003		1.00 25.73		N
	ATOM		N							D	C
10	ATOM	7425	CA	LEU	290		-89.423		1.00 27.49	D	
10	ATOM	7426	CB	LEU	290		-90.873		1.00 30.25	D	C
	MOTA	7427	CG	LEU	290		-91.055		1.00 31.14	D	C C
	MOTA	7428	CD1		290		-92.519		1.00 32.31	D	
	MOTA	7429		LEU	290		-90.217		1.00 32.28	D	C
45	MOTA	7430	C	LEU	290		-89.265		1.00 26.79	D	C
15	ATOM	7431	0	LEU	290		-88.847		1.00 26.47	D	0
	MOTA	7432	N	THR	291		-89.596		1.00 25.93	D	N
	MOTA	7433	CA	THR	291		-89.466		1.00 25.58	D	C
	ATOM	7434	CB	THR	291		-89.994		1.00 25.08	D	C
20	ATOM	7435		THR	291		-91.361		1.00 24.28	D	0
20	ATOM	7436	CG2	THR	291		-89.905		1.00 24.63	D	C
	ATOM	7437	C	THR	291		-88.003		1.00 25.32	D	C
	MOTA	7438	0	THR	291		-87.702		1.00 25.11	D	0
	MOTA	7439	N	LEU	292		-87.097		1.00 25.55	D	N
25	ATOM	7440	CA	LEU	292		-85.668		1.00 26.33	D	С
25	MOTA	7441	CB	LEU	292		-84.857		1.00 25.09	D	C
	MOTA	7442	CG	LEU	292		-83.329		1.00 24.67	D	C
	MOTA	7443		LEU	292		-82.825		1.00 23.71	D	C
	MOTA	7444		LEU	292		-82.682		1.00 23.44	D	С
30	ATOM	7445	С	LEU	292		-85.217		1.00 27.35	D	C
30	ATOM	7446	0	LEU	292		-84.492		1.00 26.47	D	0
	ATOM	7447	N	GLN	293		-85.638		1.00 29:48	D	N
	ATOM	7448	CA	GLN	293			131.928	1.00 32.08	D	С
	MOTA	7449	CB	GLN	293			131.301	1.00 34.17	D	С
25	ATOM	7450	CG	GLN	293			131.910	1.00 37.03	D	С
35	MOTA	7451	CD	GLN	293 293			131.347 130.129	1.00 39.59	D	C
	ATOM	7452	NE2	GLN					1.00 41.51	D	0
	ATOM	7453			293			132.232	1.00 40.22 1.00 32.47	D	N
	ATOM	7454	C O	GLN	293 293			131.198 130.489		D D	C O
40	MOTA	7455		GLN					1.00 31.68		
40	MOTA	7456	N CA	SER	294 294			131.386 130.760	1.00 33.78 1.00 35.43	D D	N C
	ATOM ATOM	7457 7458	CB	SER SER	294			130.760	1.00 35.43	D	C
		7459	OG	SER	294			130.763	1.00 33.37	_	_
	ATOM ATOM	7460	C	SER	294			131.130	1.00 35.13	D D	0 C
45	ATOM	7461	o	SER	294			130.282	1.00 36.34	D	0
40	ATOM	7462	N	TYR	295			130.202	1.00 35.34	D	
	MOTA	7463	CA	TYR	295			132.830	1.00 35.67	D	N C
	MOTA	7464	CB	TYR	295			134.350	1.00 35.73	D	c
	MOTA	7465	CG	TYR	295			134.855	1.00 33.73	D	C
50	ATOM	7466		TYR	295			134.827	1.00 34.72	D	c
50	ATOM	7467		TYR	295			135.226	1.00 34.72	D	c
	MOTA	7468	CD2		295			135.220	1.00 34.93	D	c
	MOTA	7469	CE2		295			135.706	1.00 33.18	D	
	ATOM	7470	CEZ	TYR	295			135.662	1.00 34.88	D	C C
55	ATOM	7471	OH	TYR	295			136.034	1.00 35.17	D	0
55	ATOM	7472	C	TYR	295			130.034	1.00 35.12	D	C
	ATOM	7473	o	TYR	295			132.119	1.00 37.83		0
	ATOM	7473		ILE	296			132.040	1.00 37.80		N
	AION	1414	TA.	1112	230	-12.111	-03.010	132.040	1.00 33.30	U	1/4

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		7475	-		206	10 745	00 514	424 272	1 00 11 00	_	_
	ATOM	7475	CA	ILE	296		-82.514		1.00 41.08	D	C
	ATOM	7476	CB	ILE	296		-81.833		1.00 40.29	D	C
	ATOM	7477	CG2		296		-80.591		1.00 39.62	D	C
5	ATOM	7478	CG1		296		-81.446		1.00 39.58	D	C
3	MOTA	7479	CD1		296		-80.836		1.00 37.65	D	C
	MOTA	7480	С	ILE	296		-82.651		1.00 42.83	D	С
	MOTA	7481	0	ILE	296		-81.844		1.00 42.30	D	0
	MOTA	7482	N	LYS	297		-83.673		1.00 45.50	D	N
40	MOTA	7483	CA	LYS	297		-83.929		1.00 48.44	D	С
10	MOTA	7484	СВ	LYS	297		-85.114		1.00 48.12	D	С
	ATOM	7485	CG	LYS	297		-84.861		1.00 48.26	D	С
	MOTA	7486	CD	LYS	297		-86.096		1.00 49.23	D	С
	MOTA	7487	CE	LYS	297		-85.891		1.00 50.24	D	С
. –	MOTA	7488	NZ	LYS	297		-87.102		1.00 50.73	D	N
15	MOTA	7489	C	LYS	297		-84.245		1.00 50.60	D	С
	MOTA	7490	0	LYS	297	-14.903	-83.690	126.707	1.00 50.62	D	0
	MOTA	7491	N	GLY	298		-85.151		1.00 53.71	D	N
	MOTA	7492	CA	GLY	298			128.300	1.00 57.58	D	C
	MOTA	7493	C	GLY	298	-17.093	-84.372	128.665	1.00 60.34	D	С
20	MOTA	7494	0	GLY	298	-17.583	-83.664	127.794	1.00 60.81	D	0
	MOTA	7495	N	GLN	299	-17.308	-84.180	129.960	1.00 63.65	D	N
	ATOM	7496	CA	GLN	299	-18.163	-83.108	130.453	1.00 66.89	D	С
	MOTA	7497	CB	GLN	299	-18.154	-83.099	131.988	1.00 67.10	D	С
	MOTA	7498	CG	GLN	299	-19.240	-82.244	132.644	1.00 68.11	D	С
25	MOTA	7499	CD	GLN	299	-19.157	-80.773	132.268	1.00 68.71	D	С
	MOTA	7500	OE1	GLN	299	-18.097	-80.153	132.367	1.00 69.05	D	0
	ATOM	7501	NE2	GLN	299	-20.283	-80.205	131.844	1.00 69.38	D	N
	ATOM	7502	С	GLN	299	-17.716	-81.745	129.926	1.00 68.91	D	С
	MOTA	7503	0	GLN	299	-16.739	-81.171	130.411	1.00 69.02	D	0
30	MOTA	7504	N	GLN	300	-18.440	-81.239	128.931	1.00 71.20	D	N
	MOTA	7505	CA	GLN	300			128.336	1.00 73.67	D	C
	MOTA	7506	СВ	GLN	300			127.836	1.00 74.21	D	C
	MOTA	7507	CG	GLN	300			126.939	1.00 75.16	D	C
	MOTA	7508	CD	GLN	300			125.951	1.00 75.58	D	C
35	ATOM	7509	OE1		300			126.340	1.00 75.70	D	ō
	ATOM	7510	NE2		300			124.662	1.00 75.83	D	N
	ATOM	7511	С	GLN	300			127.203	1.00 74.94	D	C
	ATOM	7512	Ö	GLN	300			127.454	1.00 75.30	D	ō
	ATOM	7513	Ŋ	ARG	301			125.972	1.00 76.23	D	N
40	ATOM	7514	CA	ARG	301			124.775	1.00 77.42	D	C
	MOTA	7515	СВ	ARG	301			124.734	1.00 77.89	D	c
	ATOM	7516	CG	ARG	301			123.354	1.00 78.74	D	c
	ATOM	7517	CD	ARG	301			122.388	1.00 79.60	D	c
	MOTA	7518	NE	ARG	301			122.053	1.00 79.99	D	N
45	ATOM	7519	CZ	ARG	301			121.181	1.00 80.24	D	C
40	ATOM	7520		ARG	301			120.529	1.00 80.24	D	N
	ATOM	7521		ARG	301			120.329	1.00 80.28		
		7521	C							D	N
	MOTA			ARG	301			2 124.705 123.624	1.00 77.75	D	C
50	MOTA	7523	0	ARG	301				1.00 78.10	D	0
50	ATOM	7524	N	ARG	302			125.868	1.00 77.90	D	N
	ATOM	7525	CA	ARG	302			125.980	1.00 77.67	D	C
	ATOM	7526	CB	ARG	302			126.965	1.00 78.48	D	C
	ATOM	7527		ARG	302			126.880	1.00 79.56	D	
	MOTA	7528	CD	ARG	302			125.564	1.00 80.20	D	
55	ATOM	7529		ARG	302			7 125.360	1.00 81.19	D	N
	ATOM	7530		ARG	302			7 124.499	1.00 81.77	D	
	ATOM	7531		L ARG	302			2 123.749	1.00 81.87	D	
	ATOM	7532	NH:	2 ARG	302	-25.779	-78.720	124.386	1.00 81.98	D	N

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	MOTA	7533	С	ARG	302	-18.911			1.00 76.83	D	С
	MOTA	7534	0	ARG	302		-74.206		1.00 77.05	D	0
	MOTA	7535	N	PRO	303	-18.459	-75.423	127.711	1.00 75.72	D	N
_	MOTA	7536	CD	PRO	303	-18.978	-76.402	128.687	1.00 75.41	D	С
5	MOTA	7537	CA	PRO	303	-17.215	-74.820	128.201	1.00 74.46	D	С
	MOTA	7538	CB	PRO	303	-16.951	-75.576	129.505	1.00 74.67	D	C
	ATOM	7539	CG	PRO	303	-18.322	-75.955	129.968	1.00 75.17	D	C
	MOTA	7540	С	PRO	303	-16.100			1.00 73.06	D	С
	MOTA	7541	0	PRO	303	-15.406	-74.104	126.783	1.00 73.11	D	0
10	MOTA	7542	N	ARG	304	-15.962			1.00 71.01	D	N
	MOTA	7543	CA	ARG	304	-14.947			1.00 68.70	D	С
	MOTA	7544	CB	ARG	304	-15.518			1.00 69.88	D	C
	MOTA	7545	CG	ARG	304	-14.832			1.00 71.21	D	C
	MOTA	7546	CD	ARG	304	-13.434	-77.252	123.009	1.00 72.37	D	С
15	MOTA	7547	NE	ARG	304			122.386	1.00 73.31	D	N
	MOTA	7548	$\mathbf{cz}$	ARG	304			121.254	1.00 73.71	D	C
	ATOM	7549		ARG	304			120.578	1.00 74.08	D	N
	MOTA	7550		ARG	304			120.792	1.00 73.93	D	N
	MOTA	7551	С	ARG	304			125.869	1.00 66.18	D	C
20	MOTA	7552	0	ARG	304			125.054	1.00 66.00	D	0
	MOTA	7553	N	ASP	305			126.884	1.00 63.00	D	N
	MOTA	7554	CA	ASP	305			127.091	1.00 59.34	D	С
	MOTA	7555	CB	ASP	305			128.414	1.00 59.31	D	C
0.5	MOTA	7556	CG	ASP	305			128.703	1.00 59.38	D	C
25	MOTA	7557		ASP	305			127.766	1.00 59.54	D	0
	MOTA	7558		ASP	305			129.861	1.00 59.45	D	0
	ATOM	7559	С	ASP	305			127.100	1.00 56.66	D	С
	MOTA	7560	0	ASP	305			128.036	1.00 56.22	D	0
00	MOTA	7561	Ŋ	ARG	306			126.042	1.00 52.93	D	N
30	MOTA	7562	CA	ARG	306			125.926	1.00 49.32	D	C
	ATOM	7563	СВ	ARG	306			124.487	1.00 50.93	D	C
	MOTA	7564	CG	ARG	306			123.481	1.00 52.84	D	С
	MOTA	7565	CD	ARG	306			122.147	1.00 54.60	D	С
25	ATOM	7566	NE	ARG	306			121.500	1.00 56.43	D	N
35	ATOM	7567	CZ	ARG	306			120.917	1.00 57.34	D	С
	MOTA	7568		ARG	306			120.887	1.00 58.29	D	N
	ATOM	7569		ARG	306			120.367	1.00 57.94	D	N
	ATOM	7570	C	ARG	306			126.879	1.00 45.59	D	С
40	MOTA	7571	0	ARG	306			127.086	1.00 45.21	D	0
40	MOTA	7572	N	PHE	307			127.456	1.00 41.34	D	N
	MOTA	7573	CA	PHE	307			128.378	1.00 37.64	D	C
	MOTA	7574	CB	PHE	307			128.244	1.00 37.62	D	С
	MOTA	7575	CG	PHE	307			126.876	1.00 38.89	D	C
45	MOTA	7576		PHE	307			125.872	1.00 38.94	D	C
43	ATOM	7577 7578		PHE	307 307			126.580 124.591	1.00 38.30	D	C
	ATOM	7579		PHE	307				1.00 38.89	D	С
	ATOM ATOM	7580	CEZ	PHE	307			125.305 124.309	1.00 38.37	D	C
	ATOM	7581	C	PHE	307			124.309	1.00 38.57	D	C
50	MOTA	7582	o	PHE	307			130.692	1.00 34.77	D	C
50	ATOM	7583	N	LEU	308			130.692	1.00 33.68	D	0
	ATOM	7584	CA	LEU	308			130.174	1.00 31.44 1.00 28.34	D	N
	ATOM	7585	CB	LEU	308			131.558		D	C
	ATOM	7586	CG	LEU	308			131.625	1.00 27.84 1.00 28.71	D	C
55	ATOM	7587		LEU	308			. 132.793	1.00 28.71	D	С
55	MOTA	7588		LEU	308			3 134.081	1.00 29.15	D D	C
	ATOM	7589	C	LEU	308			2 132.311	1.00 26.96	D	C
	ATOM	7590	0	LEU	308			132.311	1.00 26.41	D	С
	AIUH	, 550	9	<b>U</b> EQ	200	0.724	- 10.191	133.343	1.00 24.44	ע	0

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	MOTA	7591	N	TYR	309	-7.079 -78.385 131.810 1.00 24.84 D N	
	MOTA	7592	CA	TYR	309	-6.199 -79.333 132.489 1.00 24.89 D (	
	MOTA	7593	CB	TYR	309	-6.107 -80.634 131.691 1.00 24.35 D	
_	MOTA	7594	CG	TYR	309	-5.286 -81.708 132.373 1.00 25.38 D	
5	MOTA	7595	CD1		309	-5.667 -82.214 133.617 1.00 24.51 D	
	MOTA	7596		TYR	309	-4.922 -83.203 134.254 1.00 24.34 D	
	MOTA	7597		TYR	309	-4.127 -82.220 131.778 1.00 24.33 D	
	ATOM	7598	CE2	TYR	309		2
10	MOTA	7599	CZ	TYR	309		2
10	MOTA	7600	OH	TYR	309		2
	MOTA	7601	C	TYR	309		2
	MOTA	7602	0	TYR	309		0
	MOTA	7603	N	ALA	310		N
4 =	MOTA	7604	CA	ALA	310		0
15	MOTA	7605	CB	ALA	310		C
	MOTA	7606	C	ALA	310		C
	MOTA	7607	0	ALA	310		0
	MOTA	7608	N	LYS	311		N
00	MOTA	7609	CA	LYS	311		С
20	MOTA	7610	CB	LYS	311		С
	MOTA	7611	CG	LYS	311		С
	ATOM	7612	CD	LYS	311		С
	MOTA	7613	CE	LYS	311		С
0.5	MOTA	7614	NZ	LYS	311		N
25	MOTA	7615	С	LYS	311		С
	MOTA	7616	0	LYS	311		0
	MOTA	7617	N	LEU	312		N
	MOTA	7618	CA	LEU	312		С
	ATOM	7619	CB	LEU	312		С
30	MOTA	7620	CG	LEU	312		С
	MOTA	7621		LEU	312		C
	MOTA	7622		LEU	312		С
	MOTA	7623	С	LEU	312		C
	MOTA	7624	0	LEU	312		0
35	MOTA	7625	N	LEU	313		N
	ATOM	7626	CA	LEU	313		C
	MOTA	7627	CB	LEU	313		С
	MOTA	7628	CG	LEU	313		C
4.0	MOTA	7629		LEU	313		С
40	MOTA	7630		LEU	313		C
	MOTA	7631	С	LEU	313		С
	MOTA	7632	0	LEU	313		0
	MOTA	7633	N	GLY	314		N
4.5	MOTA	7634	CA	GLY	314		С
45	MOTA	7635	С	GLY	314		С
	MOTA	7636	0	GLY	314		0
	MOTA	7637	N	LEU	315		N
	MOTA	7638	CA	LEU	315	-2.182 -74.554 139.986 1.00 24.19 D	С
	MOTA	7639	CB	LEU	315	-3.706 -74.402 139.979 1.00 24.08 D	С
50	MOTA	7640	CG	LEU	315	-4.171 -73.057 139.404 1.00 25.31 D	С
	MOTA	7641		LEU	315	-5.668 -73.072 139.159 1.00 24.47 D	С
	MOTA	7642		LEU	315	-3.795 -71.932 140.383 1.00 24.52 D	С
	MOTA	7643	С	LEU	315	-1.741 -75.547 141.055 1.00 23.53 D	C
	MOTA	7644	0	LEU	315	-1.569 -75.169 142.213 1.00 23.85 D	0
55	MOTA	7645	N	LEU	316	-1.552 -76.806 140.667 1.00 23.29 D	N
	MOTA	7646	CA	LEU	316	-1.095 -77.834 141.605 1.00 23.36 D	C
	ATOM	7647	CB	LEU	316	-1.196 -79.222 140.982 1.00 24.00 D	С
	MOTA	7648	CG	LEU	316	-2.589 -79.846 141.012 1.00 26.29 D	С

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	7.mov	7640	CD1	TTOLY	21.0	2 522	01 247	140 416	1 00 05 07	-	^
	MOTA	7649 7650	CD1 CD2		316		-81.247 -79.904		1.00 25.87	D	C
	MOTA				316				1.00 25.54	D	С
	MOTA	7651		LEU	316		-77.572		1.00 22.98	D	C
5	MOTA	7652		LEU	316		-77.776		1.00 22.32	D	0
5	MOTA	7653	N	ALA	317		-77.134		1.00 22.45	D	N
	ATOM	7654	CA	ALA	317		-76.815		1.00 22.43	D	C
	ATOM	7655	СВ	ALA	317		-76.602		1.00 21.73	D	С
	ATOM	7656	C	ALA	317		-75.543		1.00 22.46	D	С
40	MOTA	7657	0	ALA	317		-75.435		1.00 21.94	D	0
10	MOTA	7658	N	GLU	318		-74.581		1.00 22.48	D	N
	MOTA	7659	CA	GLU	318		-73.353		1.00 23.87	D	С
	MOTA	7660	CB	GLU	318		-72.364		1.00 25.70	D	С
	ATOM	7661	CG	GLU	318		-71.114		1.00 31.16	D	С
4.5	MOTA	7662	CD	GLU	318		-69.941		1.00 34.23	D	С
15	MOTA	7663		GLU	318			141.761	1.00 36.85	D	0
	MOTA	7664			318			142.164	1.00 37.30	D	0
	MOTA	7665	C	GLU	318			144.144	1.00 23.13	D	C
	MOTA	7666	0	GLU	318			145.095	1.00 21.66	D	0
00	MOTA	7667	N	LEU	319			144.299	1.00 22.62	D	N
20	ATOM	7668	CA	LEU	319			145.623	1.00 22.59	D	C
	ATOM	7669	CB	LEU	319			145.491	1.00 22.32	D	C
	ATOM	7670	CG	LEU	319			146.743	1.00 22.65	D	С
	MOTA	7671		LEU	319			147.601	1.00 19.98	D	С
0.5	MOTA	7672		LEU	319			146.314	1.00 21.48	D	С
25	MOTA	7673	C	LEU	319			146.349	1.00 22.54	D	С
	MOTA	7674	0	LEU	319			147.560	1.00 21.29	D	0
	MOTA	7675	N	ARG	320			145.597	1.00 22.22	D	N
	MOTA	7676	CA	ARG	320			146.163	1.00 23.46	D	С
00	MOTA	7677	CB	ARG	320			145.100	1.00 25.91	D	С
30	ATOM	7678	CG	ARG	320			145.639	1.00 29.09	D	С
	MOTA	7679	CD	ARG	320			146.542	1.00 32.17	D	С
	MOTA	7680	NE	ARG	320			147.039	1.00 35.46	D	N
	ATOM	7681	CZ	ARG	320			146.590	1.00 35.28	D	С
0.5	MOTA	7682		ARG	320			145.619	1.00 35.19	D	N
35	MOTA	7683		ARG	320			147.120	1.00 36.23	D	N
	ATOM	7684	С	ARG	320			146.675	1.00 22.95	D	С
	MOTA	7685	0	ARG	320			147.758	1.00 22.89	D	0
	ATOM	7686	N	SER	321			145.893	1.00 22.56	D	N
40	ATOM	7687	CA	SER	321			146.302	1.00 23.16	D	C
40	MOTA	7688	CB	SER	321			145.210	1.00 24.44	D	С
	MOTA	7689	OG	SER	321	5.696		144.004	1.00 29.71	D	0
	MOTA	7690	C	SER	321			147.578	1.00 21.76	D	C
	MOTA	7691	0	SER	321			148.477	1.00 21.35	D	0
4 500	MOTA	7692	N	ILE	322			147.648	1.00 20.52	D	N
45	MOTA	7693	CA	ILE	322			148.825	1.00 20.27	D	C
	MOTA	7694	CB	ILE	322			148.611	1.00 19.98	D	С
	MOTA	7695	CG2		322			149.941	1.00 20.29	D	С
	MOTA	7696		ILE	322			147.601	1.00 20.19	D	C
<b>50</b>	MOTA	7697	CD1		322			147.197	1.00 21.36	D	С
50	MOTA	7698	С	ILE	322			150.054	1.00 19.11	D	С
	MOTA	7699	0	ILE	322			151.120	1.00 17.80	D	0
	MOTA	7700	N	ASN	323			149.888	1.00 20.09	D	N
	MOTA	7701	CA	ASN	323			150.972	1.00 20.89	D	C
EC	MOTA	7702	CB	ASN	323			150.449	1.00 23.19	D	С
55	ATOM	7703	CG	ASN	323			151.556	1.00 26.69	D	С
	ATOM	7704		. ASN	323			151.453	1.00 28.16	D	0
	ATOM	7705		ASN	323			152.602	1.00 26.88	D	N
	MOTA	7706	С	ASN	323	4.398	-75.369	151.511	1.00 20.52	D	C

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	3 moss	7707	•	ASN	323	4.609 -75.346 152.718 1.00 18.86 D	_
	MOTA		0		324		
	MOTA	7708	N	GLU			
	MOTA	7709	CA	GLU	324	6.763 -74.997 151.009 1.00 22.21 D	
_	MOTA	7710	CB	GLU	324	7.689 -75.077 149.794 1.00 23.95 D	
5	MOTA	7711	CG	GLU	324	8.155 -76.472 149.463 1.00 27.94 D	
	MOTA	7712	CD	GLU	324	8.511 -76.621 148.003 1.00 31.05 D	
	MOTA	7713	OE1		324	9.069 -75.663 147.424 1.00 33.20 D	
	MOTA	7714	OE2	GLU	324	8.238 -77.702 147.435 1.00 32.71 D	
	ATOM	7715	С	GLU	324	6.938 -73.630 151.671 1.00 21.32 D	С
10	MOTA	7716	0	GLU	324	7.694 -73.490 152.633 1.00 20.59 D	0
	MOTA	7717	N	ALA	325	6.256 -72.619 151.140 1.00 20.62 D	N
	ATOM	7718	CA	ALA	325	6.339 -71.281 151.707 1.00 20.95 D	C
	MOTA	7719	CB	ALA	325	5.551 -70.298 150.853 1.00 20.62 D	
	MOTA	7720	С	ALA	325	5.795 -71.317 153.139 1.00 21.67 D	
15	MOTA	7721	0	ALA	325	6.271 -70.587 154.002 1.00 23.20 D	
. •	MOTA	7722	N	TYR	326	4.802 -72.164 153.396 1.00 21.72 D	
	MOTA	7723	CA	TYR	326	4.259 -72.285 154.747 1.00 22.87 p	
	ATOM	7724	СВ	TYR	326	3.150 -73.337 154.798 1.00 21.37 E	
	ATOM	7725	CG	TYR	326	1.745 -72.783 154.727 1.00 20.49	
20	ATOM	7726		TYR	326		
20		7727					
	ATOM		CE1	TYR	326	-0.074 -71.498 155.701 1.00 18.19 I	
	MOTA	7728	CD2	TYR	326	0.889 -73.145 153.689 1.00 19.15 I	
	MOTA	7729	CE2	TYR	326	-0.422 -72.707 153.652 1.00 17.68 I	
05	MOTA	7730	CZ	TYR	326	-0.901 -71.888 154.658 1.00 18.08 I	
25	MOTA	7731	ОН	TYR	326	-2.220 -71.494 154.626 1.00 16.91 I	
	MOTA	7732	С	TYR	326	5.392 -72.697 155.688 1.00 23.78 I	
	MOTA	7733	0	TYR	326	5.551 -72.135 156.775 1.00 22.86 I	
	MOTA	7734	N	GLY	327	6.173 -73.687 155.263 1.00 24.36 I	
	MOTA	7735	CA	GLY	327	7.298 -74.147 156.064 1.00 25.31 I	C
30	MOTA	7736	С	GLY	327	8.252 -73.007 156.385 1.00 26.49 I	C
	MOTA	7737	0	GLY	327	8.747 -72.901 157.504 1.00 26.84 I	0
	MOTA	7738	N	TYR	328	8.518 -72.150 155.406 1.00 27.41 I	N C
	MOTA	7739	CA	TYR	328	9.399 -71.006 155.623 1.00 28.77 I	ОС
	MOTA	7740	СВ	TYR	328	9.628 -70.263 154.306 1.00 30.90 I	ОС
35	ATOM	7741	CG	TYR	328		C
	ATOM	7742		TYR	328		o c
	ATOM	7743	CE1		328		o c
	ATOM	7744		TYR	328		o c
	MOTA	7745	CE2		328		o c
40	ATOM	7746	CZ	TYR	328		D C
40	ATOM	7747	OH	TYR	328		0 0
	MOTA	7748	C	TYR	328		D C
	ATOM	7749	0	TYR	328		D 0
15	ATOM	7750	N	GLN	329		D N
45	MOTA	7751	CA	GLN	329		D C
	MOTA	7752	CB	GLN	329		D C
	MOTA	7753	CG	GLN	329		D C
	MOTA	7754	CD	GLN	329		D C
	ATOM	7755		GLN	329		D O
50	MOTA	7756		GLN	329		D M
	ATOM	7757	С	GLN	329		D C
	MOTA	7758	0	GLN	329	7.161 -68.624 159.761 1.00 27.55	D O
	ATOM	7759	N	ILE	330	6.659 -70.679 159.024 1.00 29.84	D N
	ATOM	7760	CA	ILE	330		D C
55	ATOM	7761	CB	ILE	330		D C
	ATOM	7762	CG2		330		D C
	ATOM	7763		ILE	330		D C
	ATOM	7764		ILE	330		D C
	1.1011	,,,,,,				/x>3 x33./14 x.00 32.0/	

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			_				_
	MOTA	7765	С	ILE	330	8.081 -71.215 161.017 1.00 33.17 D	
	MOTA	7766	0	ILE	330	8.180 -71.044 162.233 1.00 32.52 D	0
	MOTA	7767	N	GLN	331	9.142 -71.358 160.229 1.00 34.61 D	
	MOTA	7768	CA	GLN	331	10.489 -71.323 160.790 1.00 36.32 E	C
5	MOTA	7769	CB	GLN	331	11.462 -72.086 159.885 1.00 37.97 D	C
	MOTA	7770	CG	GLN	331	11.516 -73.581 160.175 1.00 41.59	C
	MOTA	7771	CD	GLN	331	11.035 -74.429 159.014 1.00 44.09 [	
	ATOM	7772		GLN	331	11.678 -74.490 157.958 1.00 45.41	
	MOTA	7773	NE2	GLN	331	9.894 -75.091 159.201 1.00 45.77 I	
10	MOTA	7774	C	GLN	331	11.033 -69.922 161.055 1.00 36.37 I	
10	ATOM	7775	Ö	GLN	331	11.827 -69.732 161.973 1.00 36.85 E	
	ATOM	7776	Ŋ	HIS	332	10.590 -68.946 160.268 1.00 36.20 I	
		7777	CA		332		
	MOTA	7778		HIS			
15	MOTA		CB	HIS	332	11.273 -66.977 159.006 1.00 38.26 I	
15	MOTA	7779	CG	HIS	332	12.444 -67.562 158.281 1.00 40.86 I	
	MOTA	7780		HIS	332	12.602 -68.754 157.656 1.00 41.83 I	
	MOTA	7781		HIS	332	13.659 -66.916 158.186 1.00 41.67 I	
	MOTA	7782		HIS	332	14.514 -67.684 157.535 1.00 42.92 I	
	MOTA	7783		HIS	332		ИС
20	MOTA	7784	С	HIS	332		C
	MOTA	7785	0	HIS	332	10.631 -65.502 161.523 1.00 34.73 I	0
	MOTA	7786	N	ILE	333	8.986 -67.025 161.577 1.00 33.52 I	N C
	MOTA	7787	CA	ILE	333	8.103 -66.154 162.343 1.00 31.90	o C
	MOTA	7788	CB	ILE	333	6.845 -65.815 161.524 1.00 32.25 1	ОС
25	MOTA	7789	CG2	ILE	333		о с
	ATOM	7790	CG1		333		D C
	ATOM	7791		ILE	333		D C
	ATOM	7792	C	ILE	333		D C
	ATOM	7793	ŏ	ILE	333		0 0
30	MOTA	7794	N	GLN	334		D N
50	ATOM	7795	CA	GLN	334		D C
	ATOM	7796	CB	GLN	334		D C
		7797	CG	GLN	334		
	MOTA						
25	MOTA	7798	CD	GLN	334		D C
35	MOTA	7799	OE1	-	334		D 0
	MOTA	7800	NE2		334		D N
	ATOM	7801	C	GLN	334		D C
	MOTA	7802	0	GLN	334		D O
4.0	MOTA	7803	N	GLY	335		D M
40	MOTA	7804	CA	GLY	335		D C
	MOTA	7805	С	GLY	335		D C
	MOTA	7806	0	GLY	335		D O
	ATOM	7807	N	LEU	336		D N
	MOTA	7808	CA	LEU	336	3.066 -69.005 163.855 1.00 23.83	D C
45	MOTA	7809	CB	LEU	336	3.673 -68.812 162.465 1.00 24.18	D C
	ATOM	7810	CG	LEU	336	3.504 -67.489 161.718 1.00 25.08	D C
	MOTA	7811	CD1	LEU	336	4.072 -67.659 160.312 1.00 25.55	D C
	MOTA	7812		LEU	336		D C
	MOTA	7813	C	LEU	336		D C
50	MOTA	7814	Ō	LEU	336		D O
	ATOM	7815	N	SER	337		D N
	MOTA	7816	CA	SER	337		D C
	ATOM	7817	CB	SER	337		D C
	ATOM	7818	OG	SER	337		D O
55			C		33 <i>7</i> 337		
J	ATOM	7819		SER			_
	MOTA	7820	0	SER	337		D 0
	ATOM	7821	N	ALA	338		D N
	MOTA	7822	CA	ALA	338	1.138 -72.272 167.556 1.00 26.59	D C

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	MOTA	7823	CB	ALA	338		-71.032		1.00 26.85	D	С
	MOTA	7824	С	ALA	338	-0.145	-72.646	166.814	1.00 27.29	D	С
	ATOM	7825	0	ALA	338	-1.029	~73.267	167.390	1.00 27.76	D	0
_	MOTA	7826	N	MET	339	-0.252	-72.262	165.545	1.00 27.52	D	N
5	MOTA	7827	CA	MET	339	-1.438	-72.576	164.760	1.00 29.68	D	С
	MOTA	7828	CB	MET	339		~71.410		1.00 26.75	D	С
	MOTA	7829	CG	MET	339	-2.256	-70.179	164.581	1.00 24.80	D	С
	MOTA	7830	SD	MET	339	-2.381	-68.721	163.556	1.00 21.82	D	S
	ATOM	7831	CE	MET	339	-0.670	-68.330	163.361	1.00 22.13	D	C
10	ATOM	7832	С	MET	339		-73.884		1.00 32.16	D	С
	MOTA	7833	0	MET	339	-2.268	-74.263	163.271	1.00 32.08	D	0
	MOTA	7834	N	MET	340	-0.204	-74.574	164.087	1.00 36.22	D	N
	ATOM	7835	CA	MET	340	-0.037	-75.858	163.409	1.00 41.37	D	С
	MOTA	7836	CB	MET	340	1.340	-76.439	163.713	1.00 42.92	D	С
15	MOTA	7837	CG	MET	340	2.436	-75.849	162.858	1.00 45.73	D	С
	MOTA	7838	SD	MET	340	2.720	-76.818	161.366	1.00 51.37	D	s
	MOTA	7839	CE	MET	340	1.237	-76.539	160.417	1.00 49.14	D	С
	MOTA	7840	С	MET	340	-1.136	-76.798	163.904	1.00 43.73	D	С
	ATOM	7841	0	MET	340	-1.386	-76.894	165.103	1.00 44.25	D	0
20	MOTA	7842	N	PRO	341	-1.799	-77.508	162.980	1.00 46.50	D	N
	MOTA	7843	CD	PRO	341			161.589	1.00 47.23	D	С
	MOTA	7844	CA	PRO	341			163.250	1.00 49.23	D	С
	MOTA	7845	СВ	PRO	341			161.929	1.00 48.42	D	C
	MOTA	7846	CG	PRO	341	-1.655	-79.094	161.315	1.00 47.95	D	С
25	ATOM	7847	С	PRO	341			164.468	1.00 51.91	D	С
	MOTA	7848	0	PRO	341			165.591	1.00 52.98	D	Ō
	MOTA	7849	N	LEU	342			164.255	1.00 54.46	D	N
	ATOM	7850	CA	LEU	342			165.360	1.00 56.95	D	C
	ATOM	7851	CB	LEU	342			164.875	1.00 57.06	D	C
30	ATOM	7852	CG	LEU	342			164.115	1.00 57.17	D	C
	MOTA	7853	CD1	LEU	342			162.658	1.00 57.93	D	Ċ
	ATOM	7854		LEU	342			164.234	1.00 57.79	D	С
	MOTA	7855	С	LEU	342			166.081	1.00 58.67	D	c
	ATOM	7856	0	LEU	342			166.115	1.00 58.76	D	O
35	MOTA	7857	N	LEU	343			166.650	1.00 60.38	D	N
	ATOM	7858	CA	LEU	343			167.391	1.00 62.04	D	C
	MOTA	7859	СВ	LEU	343			168.415	1.00 61.76	D	C
	MOTA	7860	CG	LEU	343			169.543	1.00 61.24	D	C
	MOTA	7861	CD1	LEU	343			170.236	1.00 61.38	D	Ċ
40	ATOM	7862		LEU	343	-0.036	-80.294	170.534	1.00 61.07	D	C
	MOTA	7863	С	LEU	343			166.534	1.00 63.41	D	Č
	ATOM	7864	0	LEU	343			166.024		D	
	ATOM	7865	N	GLN	344			166.373	1.00 64.75	D	N
	MOTA	7866	CA	GLN	344	3.860	-81.256	165.616	1.00 66.45	D	С
45	ATOM	7867	СВ	GLN	344			166.143	1.00 66.30	D	Ċ
	ATOM	7868	CG	GLN	344			166.091	1.00 65.84	D	c
	ATOM	7869	CD	GLN	344			165.853	1.00 65.57	D	C
	ATOM	7870		GLN	344			166.614	1.00 65.72	D	ō
	ATOM	7871		GLN	344			164.786	1.00 65.11	D	N
50	MOTA	7872	С	GLN	344			164.104	1.00 67.47	D	C
	MOTA	7873	0	GLN	344			163.541	1.00 67.50	D	ō
	ATOM	7874	N	GLU	345			163.441	1.00 68.16	D	N
	ATOM	7875	CA	GLU	345			161.991	1.00 69.15	D	C
	ATOM	7876	СВ	GLU	345			161.293	1.00 69.28	D	c
55	ATOM	7877	CG	GLU	345			160.652	1.00 69.98	D	c
	ATOM	7878	CD	GLU	345			161.622	1.00 70.48	D	C
	ATOM	7879		GLU	345			161.345	1.00 70.72	Ď	o
	ATOM	7880		GLU	345			162.651	1.00 70.72	D	0
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	MOTA	7881	С	GLU	345	2.089 -81.717 161.585 1.00 69.56 D C	:
	MOTA	7882	0	GLU	345	2.212 -82.214 160.466 1.00 69.53 D C	)
	MOTA	7883	N	ILE	346	1.228 -82.161 162.505 1.00 69.54 D N	
_	MOTA	7884	CA	ILE	346	0.306 -83.292 162.333 1.00 70.13 D C	
5	MOTA	7885	CB	ILE	346	0.604 -84.111 161.035 1.00 69.60 D C	
	MOTA	7886		ILE	346	1.800 -85.036 161.268 1.00 69.36 D C	
	MOTA	7887		ILE	346	-0.642 -84.897 160.590 1.00 68.80 D C	7
	MOTA	7888		ILE	346	-0.930 -86.167 161.379 1.00 67.66 D C	
	MOTA	7889	C	ILE	346	0.462 -84.208 163.552 1.00 70.49 D C	
10	MOTA	7890	0	ILE	346	1.504 -84.095 164.233 1.00 71.08 D (	)
	MOTA	7891	OXT	ILE	346	-0.443 -85.029 163.815 1.00 71.20 D (	)
	TER	7892		ILE	346	D	
	MOTA	7893	0	HOH	100	1.194-100.903 157.437 1.00 32.35 S	)
	MOTA	7894	0	HOH	101	5.789-101.495 107.858 1.00 41.67 S	)
15	MOTA	7895	0	нон	102	-3.286-100.790 155.260 1.00 35.57 S	)
	MOTA	7896	0	нон	103		)
	MOTA	7897	0	нон	105		2
	MOTA	7898	0	HOH	109	7.852 -54.504 152.498 1.00 13.54 S	C
	MOTA	7899	0	HOH	110	1.851 -59.616 158.936 1.00 15.42 S	C
20	MOTA	7900	0	HOH	111	4.061 -55.748 105.937 1.00 14.89 S	C
	ATOM	7901	0	нон	112	-3.981 -55.392 158.482 1.00 12.72 S	Э
	MOTA	7902	0	HOH	113		С
	MOTA	7903	0	HOH	114	-12.236 -61.051 158.803 1.00 15.08 S	0
	MOTA	7904	0	HOH	115		0
25	MOTA	7905	0	нон	116	-18.234 -71.054 169.058 1.00 16.92 S	0
	MOTA	7906	0	HOH	117		0
	MOTA	7907	0	HOH	118		0
	ATOM	7908	0	нон	119		0
	MOTA	7909	0	нон	120		0
30	MOTA	7910	0	нон	121		0
	ATOM	7911	0	нон	122		0
	MOTA	7912	0	нон	123		0
	ATOM	7913	0	нон	124		0
	MOTA	7914	0	HOH	125		0
35	ATOM	7915	0	нон	126		0
	ATOM	7916	0	нон	127		0
	ATOM	7917	0	нон	128		0
	ATOM	7918	0	нон	129		0
	ATOM	7919	0	нон	130		O
40	ATOM	7920	0	нон	131		O
	ATOM	7921	0	нон	132		0
	MOTA	7922	0	HOH	133		0
	MOTA	7923	0	HOH	134		0
	MOTA	7924	0	HOH	135		0
45	ATOM	7925	0	HOH	137		0
	MOTA	7926	0	нон			Ō
	ATOM	7927	0	HOH	139		0
	ATOM	7928	0	нон	140		0
	ATOM	7929	0	HOH	141		0
50	ATOM	7930	0	нон	142		0
	ATOM	7931	0	нон	143		Ó
	MOTA	7932	0	нон	144		ŏ
	ATOM	7933	0	нон	145		Ō
	ATOM	7934	0	нон	146		o
55	ATOM	7935	Ō	нон	147		ŏ
	ATOM	7936	Ö	нон	148		ŏ
	MOTA	7937	Ö	нон	149		ŏ
	ATOM	7938	Ö	нон	150		ŏ
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	ATOM	7939	0	нон	151	-5.245 -53.158 95.611 1.00 17.95 S O
	MOTA	7940	0	нон	152	5.815 -59.391 152.298 1.00 19.34 S O
	MOTA	7941	0	нон	153	6.055 -55.938 171.262 1.00 19.01 S O
	MOTA	7942	0	HOH	154	9.371 -61.052 112.669 1.00 19.61 S O
5	MOTA	7943	0	нон	155	-2.818 -61.077 171.684 1.00 18.36 S O
	MOTA	7944	0	HOH	156	7.580 -55.929 112.685 1.00 20.71 S O
	MOTA	7945	0	нон	157	-1.829 -35.248 152.635 1.00 21.17 S O
	MOTA	7946	0	HOH	158	-12.998 -38.744 111.352 1.00 22.19 S O
	MOTA	7947	0	HOH	159	-10.033 -57.313 153.074 1.00 26.61 S O
10	ATOM	7948	0	нон	160	-14.613 -42.493 121.850 1.00 21.46 S O
	MOTA	7949	0	HOH	161	-0.465 -59.730 96.207 1.00 22.15 S O
	MOTA	7950	0	нон	162	-1.457 -79.928 119.289 1.00 25.62 S O
	ATOM	7951	О	НОН	163	-2.242 -34.614 155.101 1.00 24.72 S O
	MOTA	7952	0	нон	164	-2.488 -28.817 149.752 1.00 24.52 S O
15	MOTA	7953	0	HOH	165	-14.811 -78.255 132.956 1.00 26.75 S O
	MOTA	7954	0	нон	166	8.396 -73.450 91.962 1.00 24.27 S O
	MOTA	7955	0	HOH	167	2.582 -61.210 92.554 1.00 19.81 S O
	MOTA	7956	0	нон	168	0.065 -36.237 163.877 1.00 22.12 S O
	MOTA	7957	0	HOH	169	9.793 -55.410 168.376 1.00 22.34 S O
20	MOTA	7958	0	HOH	170	-8.169 -68.504 136.548 1.00 25.65 S O
	MOTA	7959	0	нон	171	-7.200 -55.767 151.754 1.00 21.47 S O
	MOTA	7960	0	нон	172	-12.681 -37.331 109.182 1.00 20.87 S O
	ATOM	7961	0	НОН	173	16.665 -83.359 109.157 1.00 22.25 S O
05	MOTA	7962	0	НОН	174	8.904 -17.589 154.270 1.00 32.11 S O
25	ATOM	7963	0	НОН	175	-9.871 -57.482 160.770 1.00 22.21 S O
	MOTA	7964	0	нон	176	10.054 -68.265 94.306 1.00 22.03 S O
	MOTA	7965	0	нон	177	-7.645 -35.523 134.429 1.00 33.35 S O
	MOTA	7966	0	нон	178	-17.960 -31.774 109.544 1.00 26.39 S O
20	MOTA	7967	0	нон	179	2.162 -42.472 136.264 1.00 25.98 S O
30	MOTA	7968	0	НОН	180	3.481 -55.715 96.425 1.00 21.71 S O
	MOTA	7969	0	нон	181	-1.946 -59.108 170.157 1.00 21.79 S O
	ATOM	7970	0	нон	182	0.579 -86.300 149.857 1.00 29.11 S O
	MOTA	7971	0	нон	183	-7.526 -45.845 127.493 1.00 22.42 S O
25	ATOM	7972	0	нон	184	-17.611 -43.737 111.356 1.00 24.87 S O
35	ATOM	7973	0	нон	185	-7.509 -69.242 100.449 1.00 23.13 S O
	ATOM	7974	0	HOH	186	-5.332 -93.514 158.577 1.00 22.53 S O
	MOTA	7975	0	HOH	187	10.436 -93.075 106.554 1.00 23.92 S O
	ATOM	7976	0	HOH	188	9.310 -52.599 171.929 1.00 27.84 S O 16.827 -47.166 149.767 1.00 22.64 S O
40	MOTA	7977	0	HOH	189	
40	MOTA	7978 7979	0	HOH	190 191	2.500 -15.666 156.796 1.00 24.25 S O 0.395 -59.578 168.198 1.00 26.75 S O
	MOTA		0	HOH		• · · · · · · · · · · ·
	ATOM ATOM	7980 7981	0	нон нон	192 193	8.107 -33.555 114.378 1.00 28.48 S O -7.319 -49.935 168.792 1.00 27.62 S O
	ATOM	7982	Ö	НОН	194	-8.208 -31.818 100.278 1.00 23.15 S O
45	ATOM	7983	Ö	нон	195	7.764 -47.435 170.758 1.00 21.29 S O
40	ATOM	7984	ő	нон	196	7.480 -88.122 101.942 1.00 26.44 S O
	ATOM	7985	Ö	нон	197	14.406 -62.102 94.383 1.00 23.98 S O
	ATOM	7986	ő	нон	198	-15.741 -40.088 122.107 1.00 26.12 S O
	ATOM	7987	ő	нон	199	-1.448 -56.070 170.098 1.00 28.03 S O
50	ATOM	7988	ő	нон	200	5.541 -47.303 95.880 1.00 28.11 S O
00	ATOM	7989	ő	нон	201	-12.575 -56.737 102.862 1.00 31.08 S O
	ATOM	7990	Ö	нон	202	1.715 -58.973 94.236 1.00 25.62 S O
	ATOM	7991	Ö	нон	203	9.480 -63.602 164.661 1.00 25.82 S O
	ATOM	7992	Ö	нон	204	-1.232 -35.010 144.845 1.00 24.54 S O
55	ATOM	7993	Ö	нон	205	7.562 -71.697 147.929 1.00 26.32 S O
	ATOM	7994	Ö	нон	207	-7.835 -53.973 94.498 1.00 22.20 S O
	ATOM	7995		нон	208	7.846 -54.379 169.964 1.00 24.84 S O
	ATOM	7996		нон	209	2.796 -35.267 111.786 1.00 27.64 S O
	0.1				_0,	

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	ATOM	7997	0	НОН	210			135.459	1.00 24.54	S	0
	MOTA	7998	0	HOH	211	-18.964			1.00 21.20	S	0
	MOTA	7999	0	нон	212	-17.685			1.00 27.48	S	0
_	MOTA	8000	0	нон	213		-89.525		1.00 28.97	S	0
5	MOTA	8001	0	нон	214			111.888	1.00 34.69	S	0
	MOTA	8002	0	нон	215	-15.068			1.00 27.99	S	0
	MOTA	8003	0	нон	216	-18.184			1.00 20.26	S	0
	MOTA	8004	0	нон	217			128.888	1.00 25.97	S	0
40	MOTA	8005	0	нон	218		-78.683	97.186	1.00 27.25	s	0
10	MOTA	8006	0	нон	219			135.217	1.00 21.04	S	0
	MOTA	8007	0	HOH	220			137.131	1.00 21.84	S	0
	ATOM	8008	0	HOH	221			155.199	1.00 29.72	S	0
	ATOM	8009	0	нон	222			164.111	1.00 23.90	S	0
	MOTA	8010	0	HOH	223			160.393	1.00 21.15	S	0
15	MOTA	8011	0	HOH	224			168.538	1.00 23.30	S	0
	MOTA	8012	0	HOH	225			169.854	1.00 25.60	S	0
	MOTA	8013	0	нон	226			110.725	1.00 25.49	S	0
	MOTA	8014	0	HOH	227			169.157	1.00 27.54	S	0
	MOTA	8015	0	HOH	228			170.186	1.00 26.01	S	0
20	MOTA	8016	0	HOH	229			163.830	1.00 24.68	S	0
	MOTA	8017	0	HOH	230			149.831	1.00 27.89	S	0
	MOTA	8018	0	HOH	231		-59.422	93.575	1.00 25.99	S	0
	MOTA	8019	0	HOH	232			127.967	1.00 28.31	S	0
	MOTA	8020	0	HOH	233		-55.469		1.00 24.15	S	0
25	ATOM	8021	0	HOH	234			159.042	1.00 24.72	S	0
	MOTA	8022	0	HOH	235			151.433	1.00 35.81	S	0
	MOTA	8023	0	HOH	236			104.435	1.00 29.28	S	0
	MOTA	8024	0	HOH	237		-60.351		1.00 22.39	S	0
	MOTA	8025	0	HOH	238			142.529	1.00 25.98	S	0
30	MOTA	8026	0	HOH	239		-54.014		1.00 28.69	S	0
	MOTA	8027	0	HOH	240			134.614	1.00 27.47	S	0
	MOTA	8028	0	HOH	241	7.889	-69.278	128.039	1.00 25.45	S	0
	MOTA	8029	0	HOH	242		-56.173		1.00 25.84	S	0
	MOTA	8030	0	HOH	243		-43.127		1.00 31.21	S	0
35	MOTA	8031	0	HOH	244			171.337	1.00 24.49	S	0
	MOTA	8032	0	нон	245			164.313	1.00 26.33	S	0
	MOTA	8033	0	HOH	246			100.804	1.00 35.04	S	0
	MOTA	8034	0	HOH	247			123.050	1.00 27.65	S	0
	MOTA	8035	0	HOH	248			144.939	1.00 33.14	S	0
40	MOTA	8036	0	HOH	249			103.311	1.00 25.83	S	0
	ATOM	8037	0	HOH	250			149.641	1.00 32.57	S	0
	MOTA	8038	0	HOH	251			153.382	1.00 27.18	S	•
	MOTA	8039	0	НОН	252			149.250	1.00 26.80	S	0
	MOTA	8040	0	HOH	253			114.494	1.00 26.39	S	0
45	ATOM	8041	0	HOH	254			89.646	1.00 30.34	S	0
	MOTA	8042	0	HOH	255			155.492	1.00 32.02	S	0
	MOTA	8043	0	HOH	256			109.375	1.00 28.54	S	0
	MOTA	8044	0	HOH	257			153.151	1.00 29.77	S	0
	MOTA	8045	0	HOH	258			143.846	1.00 33.05	S	0
50	MOTA	8046	0	HOH	259			149.403	1.00 23.81	S	0
	MOTA	8047	0	HOH	260			161.866	1.00 27.32	S	0
	ATOM	8048	0	нон	261			154.705		S	0
	MOTA	8049	0	нон	263			156.022	1.00 33.90	S	0
	ATOM	8050	0	нон	264			145.484		S	0
55	ATOM	8051	0	нон	265			138.485		S	0
	MOTA	8052	0	нон	266		-76.738			S	0
	MOTA	8053	0	HOH	267			114.770		S	0
	MOTA	8054	0	нон	268	14.544	-43.496	170.824	1.00 25.33	S	0

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	MOTA	8055	0	нон	269	-10.804	-71.018	173.076	1.00 32.63	S	0
	MOTA	8056	0	HOH	270	3.311	-62.924	146.402	1.00 29.18	S	0
	MOTA	8057	0	HOH	271	5.357	-53.638	90.691	1.00 31.79	S	0
_	MOTA	8058	0	HOH	272	5.227	-86.281	154.159	1.00 31.53	S	0
5	MOTA	8059	0	HOH	273		-19.105		1.00 29.74	S	0
	MOTA	8060	0	HOH	274	-20.468	-80.759	149.925	1.00 33.49	S	0
	MOTA	8061	0	HOH	275	-5.662	-97.233	157.623	1.00 30.30	S	0
	MOTA	8062	0	HOH	276	15.625	-37.089	150.881	1.00 35.21	S	0
	ATOM	8063	0	HOH	277	12.161	-61.189	105.768	1.00 33.96	S	0
10	ATOM	8064	0	HOH	278	3.234	-34.573	109.191	1.00 33.68	S	0
	MOTA	8065	0	HOH	279	-8.880	-47.618	151.419	1.00 28.74	S	0
	ATOM	8066	0	нон	280	-22.502			1.00 27.73	S	0
	ATOM	8067	0	нон	281		-44.170		1.00 31.74	S	0
	ATOM	8068	0	нон	282		-48.188		1.00 31.25	s	0
15	MOTA	8069	0	нон	283		-87.258		1.00 29.49	S	0
	ATOM	8070	0	нон	284			161.206	1.00 27.49	S	0
	ATOM	8071	0	нон	285		-61.174	95.346	1.00 25.49	s	0
	ATOM	8072	0	нон	286			146.840	1.00 28.55	S	Ō
	ATOM	8073	0	нон	287			161.333	1.00 30.97	S	ō
20	ATOM	8074	0	нон	288			107.050	1.00 31.85	s	ō
	ATOM	8075	0	нон	289		-65.295		1.00 31.60	Š	ō
	ATOM	8076	0	нон	290			155.699	1.00 35.00	s	ō
	ATOM	8077	0	нон	291			152.640	1.00 28.95	s	ō
	ATOM	8078	Ō	нон	292			160.774	1.00 34.06	s	ŏ
25	ATOM	8079	0	НОН	293			112.721	1.00 25.07	S	ō
	ATOM	8080	ō	нон	294			160.924	1.00 37.13	s	ō
	ATOM	8081	Ō	НОН	295			171.437	1.00 31.26	s	o
	ATOM	8082	ō	НОН	296			175.216	1.00 31.58	s	Ö
	ATOM	8083	ō	НОН	297		-55.653		1.00 25.65	s	o
30	ATOM	8084	ō	нон	298		-57.553		1.00 25.05	s	o
•	ATOM	8085	Ö	НОН	299			169.041	1.00 27.46	s	o
	ATOM	8086	Ö	нон	300		-71.750		1.00 27.40	S	o
	ATOM	8087	ō	нон	301		-50.185		1.00 35.30	S	0
	ATOM	8088	ŏ	нон	302			170.669	1.00 33.30	S	o
35	ATOM	8089	Ö	нон	303		-66.338		1.00 23.25	S	o
•	ATOM	8090	ŏ	нон	304		-67.023		1.00 35.13	S	o
	ATOM	8091	ō	нон	305			170.536	1.00 35.57	S	o
	ATOM	8092	ŏ	нон	306		-27.751		1.00 20.33	S	0
	ATOM	8093	ŏ	нон	307		-71.189		1.00 23.20	S	o
40	ATOM	8094	ŏ	нон	308		-17.923		1.00 41.32	S	o
. •	ATOM	8095	Ö	нон	309	6.943		136.885	1.00 41.20	S	o
	ATOM	8096	ŏ	нон	310			106.465	1.00 34.52	S	ő
	ATOM	8097	ő	нон	311			166.802	1.00 34.32	S	0
	ATOM	8098	ő	нон	312			123.176	1.00 29.99	S	0
45	ATOM	8099	ő	нон	313			169.043	1.00 27.78	S	0
-10	ATOM	8100	ő	нон	314			118.169	1.00 27.78	S	
	MOTA	8101	ŏ	нон	315			177.428	1.00 39.70	S	0
	ATOM	8102	Ö	НОН	316			153.276		S	0
	MOTA	8102	Ö	НОН	317			153.276	1.00 26.59		0
50	MOTA	8104		нон	318			109.924	1.00 30.90	S	0
50			0						1.00 35.70	S	0
	ATOM	8105	0	HOH	319		-57.093		1.00 27.84	S	0
	ATOM	8106	0	нон	320			141.033	1.00 34.15	S	0
	ATOM	8107	0	НОН	321			129.073	1.00 35.91	S	0
E E	MOTA	8108	0	НОН	323			108.859	1.00 40.25	S	0
55	MOTA	8109	0	НОН	324			153.806	1.00 29.58	S	0
	MOTA	8110	0	НОН	325			148.698	1.00 38.84	S	0
	MOTA	8111	0	НОН	326			151.149	1.00 33.24	S	0
	MOTA	8112	0	нон	327	13.585	-37.762	154.701	1.00 36.77	S	0

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	MOTA	8113	0	нон	328	-4.082	-52.659	145.859	1.00 35.37	s	0
	MOTA	8114	0	HOH	329	-12.214	-97.380	152.729	1.00 40.91	S	0
	MOTA	8115	0	HOH	330	21.253	-78.801	100.196	1.00 41.54	S	0
	ATOM	8116	0	HOH	331	20.349	-66.162	108.672	1.00 39.01	S	0
5	ATOM	8117	0	нон	332	14.913	-47.738	137.531	1.00 41.04	S	0
	ATOM	8118	0	нон	333	4.138	-44.506	175.935	1.00 34.58	S	0
	MOTA	8119	0	НОН	334	-23.111	-25.163	112.021	1.00 44.12	s	0
	ATOM	8120	0	нон	335	14.393	-21.278	151.249	1.00 38.52	S	0
	ATOM	8121	0	HOH	336	-7.214	-56.582	102.979	1.00 30.20	S	0
10	MOTA	8122	0	нон	337	-10.229	-62.939	99.872	1.00 30.38	S	0
	MOTA	8123	0	HOH	338	19.511	-44.397	155.298	1.00 29.41	S	0
	MOTA	8124	0	нон	339		-74.211	95.233	1.00 32.45	S	0
	ATOM	8125	0	нон	340	-3.916	-30.077	164.362	1.00 31.78	s	0
	MOTA	8126	0	нон	341		-80.459		1.00 28.04	S	0
15	ATOM	8127	0	нон	343	-14.436	-22.122	109.753	1.00 42.19	S	0
	ATOM	8128	0	нон	344			170.396	1.00 27.91	S	0
	MOTA	8129	0	нон	345		-22.558	160.678	1.00 43.95	S	0
	ATOM	8130	0	нон	346	18.274	-93.064	117.144	1.00 35.29	S	0
	ATOM	8131	0	нон	347	-8.246	-70.761	116.848	1.00 34.25	S	0
20	MOTA	8132	0	нон	348			113.302	1.00 34.77	S	0
	MOTA	8133	0	нон	349	-23.016	-83.847	145.069	1.00 31.88	S	0
	ATOM	8134	0	нон	350			168.903	1.00 37.36	S	Ō
	MOTA	8135	0	нон	351			153.017	1.00 32.52	S	0
	ATOM	8136	0	нон	352			156.021	1.00 48.54	S	ō
25	MOTA	8137	0	нон	353	6.820	-51.420	145.123	1.00 26.19	S	0
	ATOM	8138	0	нон	354			166.993	1.00 42.35	s	Ö
	ATOM	8139	0	нон	355			153.100	1.00 30.60	S	Ō
	ATOM	8140	o	нон	356			173.508	1.00 47.01	s	ō
	ATOM	8141	Ō	нон	358			128.237	1.00 31.19	S	Ō
30	ATOM	8142	0	нон	359			156.601	1.00 29.22	S	Ō
	ATOM	8143	0	нон	360			100.980	1.00 27.83	s	ō
	MOTA	8144	Ō	нон	361		-36.663		1.00 37.52	S	Ō
	ATOM	8145	O	нон	362			149.391	1.00 39.87	S	ō
	MOTA	8146	0	нон	363			164.566	1.00 29.90	s	ō
35	MOTA	8147	0	нон	364			135.114	1.00 32.66	S	Ō
	ATOM	8148	Ō	нон	365			157.785	1.00 32.62	S	ō
	ATOM	8149	Ó	нон	366			109.527	1.00 39.14	S	ŏ
	ATOM	8150	O	нон	367			159.935	1.00 30.91	S	Ō
	ATOM	8151	Ō	НОН	369			103.921	1.00 33.02	S	Ō
40	ATOM	8152	Ō	нон	370			128.744	1.00 35.96	S	ō
	ATOM	8153	ō	нон	371			122.607	1.00 39.74	s	ŏ
	ATOM	8154	0	нон	372			170.898	1.00 33.17	S	ō
	ATOM	8155	0	нон	373			157.889	1.00 47.16	S	Ö
	ATOM	8156	0	нон	374			173.306	1.00 31.81	s	ō
45	ATOM	8157	0	нон	375			156.839	1.00 43.43	S	Ö
	ATOM	8158	Ō	нон	376			147.266	1.00 33.03	S	ō
	ATOM	8159	0	нон	377			152.019	1.00 31.07	S	0
	ATOM	8160	Ö	нон	378			134.105	1.00 37.41	S	o
	ATOM	8161	Ō	нон	379			157.807	1.00 44.33	S	Ö
50	MOTA	8162	Ó	нон	380			127.555	1.00 35.49	S	Ö
	ATOM	8163	ŏ	нон	381			173.073	1.00 35.86	S	Ö
	ATOM	8164	ō	нон	383			3 108.536	1.00 37.37	s	Ö
	ATOM	8165	Ö	нон	385			129.029	1.00 41.45	s	ŏ
	ATOM	8166	ō	НОН	386			105.307	1.00 44.00	s	ŏ
55	MOTA	8167	Ö	нон	387			164.207	1.00 37.54	s	o
	ATOM	8168	Ö	нон	388			130.123	1.00 34.65	s	ŏ
	ATOM	8169	ő	нон	389			5 114.570	1.00 35.97	S	ŏ
	ATOM	8170	Ö	нон	390			5 135.885	1.00 30.52	S	ő
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	MOTA	8171	0	HOH	391	-8.453 -70.094 98.112 1.00 40.15 S O
	MOTA	8172	0	HOH	392	-4.178 -80.568 127.822 1.00 31.47 S O
	MOTA	8173	0	нон	393	-17.842 -63.215 143.635 1.00 39.38 S O
_	MOTA	8174	0	HOH	394	-19.264 -81.648 157.206 1.00 32.27 S O
5	MOTA	8175	0	нон	396	-20.956 -75.317 171.042 1.00 33.04 S O
	MOTA	8176	0	HOH	397	1.082 -45.036 161.177 1.00 36.77 S O
	MOTA	8177	0	нон	399	-16.486 -93.083 157.191 1.00 38.04 S O
	MOTA	8178	0	HOH	400	18.520 -69.299 94.579 1.00 38.46 S O
	MOTA	8179	0	HOH	401	-14.376 -47.406 162.731 1.00 43.61 S O
10	MOTA	8180	0	HOH	402	-15.725 -46.681 93.561 1.00 36.70 S O
	MOTA	8181	0	HOH	403	15.449 -93.447 107.056 1.00 42.96 S O
	ATOM	8182	0	нон	404	17.756 -34.951 160.597 1.00 38.74 S O
	MOTA	8183	0	HOH	405	-11.516 -52.980 92.984 1.00 31.69 S O
	MOTA	8184	0	HOH	406	2.904 -99.028 157.874 1.00 35.56 S O
15	MOTA	8185	0	HOH	407	3.990 -47.232 174.280 1.00 54.42 S O
	MOTA	8186	0	HOH	408	5.363 -82.052 150.581 1.00 39.03 S O
	MOTA	8187	0	HOH	409	-13.329 -21.194 112.697 1.00 38.48 S O
	MOTA	8188	0	HOH	410	-20.132 -46.572 109.024 1.00 42.34 S O
	MOTA	8189	0	HOH	411	-20.146 -71.594 142.398 1.00 48.84 S O
20	MOTA	8190	0	HOH	413	7.810 -96.588 117.307 1.00 48.54 S O
	MOTA	8191	0	HOH	414	-15.707 -39.444 112.856 1.00 30.16 S O
	ATOM	8192	0	HOH	415	-7.705 -43.981 141.677 1.00 42.54 S O
	ATOM	8193	0	нон	416	-11.327 -80.716 166.581 1.00 29.92 S O
	ATOM	8194	0	HOH	417	14.765 -15.174 132.791 1.00 46.34 S O
25	MOTA	8195	0	нон	418	12.180 -35.626 148.220 1.00 43.95 S O
	ATOM	8196	0	HOH	419	-0.687 -69.344 171.152 1.00 37.41 S O
	ATOM	8197	0	HOH	420	-11.442 -58.044 158.798 1.00 36.92 S O
	ATOM	8198	0	нон	421	12.102 -62.691 93.583 1.00 34.13 S O
	MOTA	8199	0	нон	422	-3.773 -38.085 92.262 1.00 52.29 S O
30	ATOM	8200	0	нон	423	12.749 -96.511 113.885 1.00 39.83 S O
	ATOM	8201	0	нон	424	2.752 -31.848 124.578 1.00 44.26 S O
	ATOM	8202	0	HOH	425	12.704 -41.353 112.133 1.00 44.28 S O
	MOTA	8203	0	нон	427	-10.440 -68.616 172.311 1.00 32.78 S O
	MOTA	8204	0	нон	428	-6.952 -47.164 90.926 1.00 36.61 S O
35	ATOM	8205	0	нон	429	-5.052 -57.339 148.491 1.00 39.32 S O
	ATOM	8206	0	HOH	431	-1.897 -61.286 90.062 1.00 52.91 S O
	MOTA	8207	0	HOH	432	17.295 -72.124 95.247 1.00 51.20 S O
	MOTA	8208	0	HOH	433	-8.442 -44.268 167.166 1.00 38.87 S O
	ATOM	8209	0	нон	434	17.130 -51.752 145.075 1.00 37.31 S O
40	MOTA	8210	0	НОН	435	-11.934 -41.853 89.278 1.00 35.15 S O
	MOTA	8211	0	HOH	436	-20.655 -79.110 158.422 1.00 38.37 S O
	MOTA	8212	0	HOH	437	-17.692 -58.619 152.324 1.00 39.50 S O
	ATOM	8213	0	HOH	438	-13.228 -92.072 160.458 1.00 42.10 S O
	MOTA	8214	0	HOH	439	4.172 -12.941 153.839 1.00 33.74 S O
45	ATOM	8215	0	HOH	440	-10.145 -87.720 162.303 1.00 38.35 S O
	ATOM	8216	0	HOH	442	-10.931 -61.150 93.207 1.00 45.01 S O
	MOTA	8217	0	HOH	443	-1.358 -40.495 93.770 1.00 37.78 S O
	ATOM	8218	0	HOH	446	-6.450 -68.499 173.937 1.00 37.62 S O
	ATOM	8219	0	HOH	448	24.212 -70.506 107.734 1.00 48.81 S O
50	ATOM	8220	0	HOH	449	-6.596 -43.865 93.009 1.00 32.44 S O
	ATOM	8221	0	нон	450	-7.000 -38.908 142.094 1.00 43.17 S O
	MOTA	8222	0	HOH	452	11.741 -63.298 155.435 1.00 33.39 S O
	ATOM	8223	0	HOH	453	-4.596 -14.237 111.600 1.00 45.05 S O
	ATOM	8224	0	нон	454	16.341 -40.682 142.327 1.00 37.42 S O
55	MOTA	8225	0	нон	455	1.573 -61.767 174.211 1.00 40.92 S O
	MOTA	8226	0	нон	456	-13.624 -35.519 117.025 1.00 35.47 S O
	MOTA	8227	0	нон	458	-16.126 -56.335 151.820 1.00 35.23 S O
	MOTA	8228	0	нон	459	17.663 -37.767 143.893 1.00 40.21 S O

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	MOTA	8229	0	нон	461	-12.126			1.00 53.95	S	0
	MOTA	8230	0	HOH	462		-12.385		1.00 34.36	S	0
	MOTA	8231	0	HOH	463		-61.641		1.00 42.87	S	0
_	MOTA	8232	0	HOH	464		-88.839		1.00 51.28	S	0
5	MOTA	8233	0	HOH	465		-57.821		1.00 39.77	S	0
	MOTA	8234	0	HOH	466		-70.873	125.729	1.00 43.43	S	0
	MOTA	8235	0	HOH	467	2.705	-61.794	90.011	1.00 43.72	S	0
	MOTA	8236	0	HOH	468			160.863	1.00 44.03	S	0
	MOTA	8237	0	HOH	469	2.275	-34.938	119.431	1.00 37.93	S	0
10	MOTA	8238	0	HOH	470	-3.650	-77.137	108.828	1.00 47.92	S	0
	MOTA	8239	0	HOH	472	-4.726	-64.694	95.196	1.00 35.25	S	0
	MOTA	8240	0	нон	473	-22.754	-72.240	159.906	1.00 34.93	S	0
	MOTA	8241	0	HOH	474	16.681	-66.920	120.698	1.00 34.05	S	0
	ATOM	8242	0	HOH	475	-10.065	-60.263	173.767	1.00 42.11	S	0
15	MOTA	8243	0	HOH	476	-6.797	-14.864	110.275	1.00 38.54	S	0
	MOTA	8244	0	HOH	477	16.699	-92.866	125.208	1.00 39.15	S	0
	MOTA	8245	0	HOH	478	-8.371	-55.088	161.316	1.00 43.44	S	0
	MOTA	8246	0	HOH	479	-10.360	-72.131	177.271	1.00 57.30	S	0
	ATOM	8247	0	нон	481	-10.827	-16.909	116.124	1.00 45.61	S	0
20	ATOM	8248	0	HOH	483	3.497	-28.876	114.517	1.00 38.30	S	0
	MOTA	8249	0	нон	485	16.708	-79.915	98.646	1.00 36.18	S	0
	MOTA	8250	0	нон	486	5.685	-70.865	142.719	1.00 39.47	S	0
	ATOM	8251	0	нон	487	6.377	-57.579	115.024	1.00 44.49	S	0
	ATOM	8252	0	нон	489			171.632	1.00 41.31	s	0
25	ATOM	8253	Ō	нон	490			161.598	1.00 53.30	S	O
	MOTA	8254	ō	НОН	491			129.532	1.00 41.18	S	0
	MOTA	8255	ō	нон	492			154.533	1.00 36.96	S	O
	ATOM	8256	ō	нон	493		-81.316		1.00 35.88	s	Ö
	ATOM	8257	Ö	НОН	494			121.694	1.00 33.42	s	ŏ
30	ATOM	8258	Ö	нон	495			112.568	1.00 35.91	s	ŏ
00	ATOM	8259	ŏ	нон	496			154.593	1.00 47.05	s	ŏ
	MOTA	8260	ő	нон	497			148.612	1.00 39.22	s	ŏ
	ATOM	8261	Ö	нон	498			103.261	1.00 39.63	S	ō
	MOTA	8262	ŏ	нон	499			113.518	1.00 34.44	s	ŏ
35	ATOM	8263	ő	нон	500			135.894	1.00 47.58	s	ŏ
00	ATOM	8264	ő	нон	501			115.223	1.00 39.61	s	ō
	ATOM	8265	Ö	нон	502			124.095	1.00 51.93	s	ŏ
	ATOM	8266	Ö	нон	503			140.453	1.00 46.40	s	ŏ
	ATOM	8267	ő	нон	504			134.440	1.00 46.55	s	ŏ
40	MOTA	8268	ő	нон	505		-68.377		1.00 46.90	s	ő
40	ATOM	8269	Ö	нон	506			108.740	1.00 42.45	s	ŏ
	ATOM	8270	ő	нон					1.00 42.69		ŏ
	ATOM	8271	Ö	нон	509			107.564	1.00 43.38	S	ŏ
		8272	Ö	НОН	510			155.934	1.00 48.47	S	ŏ
45	MOTA	8272	0	НОН	511			135.988	1.00 43.55	S	o
45	MOTA				512			114.704	1.00 43.33	S	
	ATOM	8274	0	HOH	512			157.603	1.00 39.28	S	0
	ATOM	8275	0	HOH							0
	MOTA	8276	0	НОН	514			7 143.859	1.00 48.19	S	0
<b>E</b> 0	ATOM	8277	0	НОН	515			9 113.699	1.00 41.10	S	0
50	ATOM	8278	0	НОН	516			162.242	1.00 53.03	S	0
	ATOM	8279	0	нон	517			8 120.083	1.00 50.51	S	0
	ATOM	8280	0	нон	518			3 158.553	1.00 34.18	S	
	MOTA	8281	0	нон	519		-65.95			S	
65	MOTA	8282		нон	520			0 158.067		S	
55	MOTA	8283		нон	521			2 107.777			
	ATOM	8284		нон	523		-57.46				
	MOTA	8285		нон	524			7 104.702			
	MOTA	8286	0	нон	525	23.766	-78.86	5 111.717	1.00 49.25	S	0

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	ATOM	8287	0	нон	526	12.535 -38.204 148.153 1.00 53.45 S O	
	ATOM	8288	0	нон	527	-6.678 -65.953 114.241 1.00 45.88 S O	
	MOTA	8289	Ö	нон	529	-21.878 -58.439 164.200 1.00 48.19 S O	
	MOTA	8290	ŏ	нон	530	-11.825 -37.756 116.383 1.00 45.97 S O	
5	MOTA	8291	o	нон	532	-11.859 -61.199 109.587 1.00 49.73 S O	
J	ATOM	8292	ŏ	нон	533	-5.953 -38.372 156.605 1.00 43.27 S O	
	ATOM	8293	ŏ	нон	535	-9.521 -45.912 129.467 1.00 40.80 S O	
	ATOM	8294	ŏ	нон	536	15.652 -33.654 161.052 1.00 39.31 S O	
	ATOM	8295	ŏ	нон	537	-11.557 -73.270 111.665 1.00 52.60 S O	
10	MOTA	8296	ŏ	нон	538	14.976 -96.434 109.861 1.00 51.75 S O	
10	ATOM	8297	0	нон	539	-6.961 -47.480 169.785 1.00 42.26 S O	
	ATOM	8298	ŏ	нон	540	-3.653 -40.089 169.579 1.00 44.31 S O	
	ATOM	8299	Ö	нон	542	4.492 -99.754 106.625 1.00 42.46 S O	
	ATOM	8300	ŏ	нон	543	-15.385 -59.003 171.024 1.00 38.14 S O	
15	ATOM	8301	Ö	нон	544	-9.565 -39.031 162.703 1.00 43.59 S O	
13	ATOM	8302	Ö	нон	545	-2.257 -14.277 104.237 1.00 51.26 S O	
	ATOM	8303	o	нон	546	-8.109 -97.785 106.812 1.00 39.90 S O	
		8304	0	нон	547	7.818 -98.465 145.193 1.00 48.51 S O	
	ATOM	8305		НОН	548		
20	MOTA		0		549		
20	ATOM	8306	0	НОН			
	MOTA	8307	0	нон	551 552		
	MOTA	8308	0	нон	552		
	MOTA	8309	0	HOH	553	10.549 -88.842 145.088 1.00 55.30 S O	
0E	MOTA	8310	0	нон	554	-19.934 -72.006 149.518 1.00 43.14 S O	
25	MOTA	8311	0	нон	555	-9.143 -35.862 147.215 1.00 44.48 S O	
	ATOM	8312	0	нон	557	11.968 -27.647 165.607 1.00 39.72 S O	
	MOTA	8313	0	НОН	558	7.500 -78.286 152.237 1.00 42.78 S O	
	ATOM	8314	0	нон	559	-20.328 -67.927 149.359 1.00 42.48 S O	
20	MOTA	8315	0	НОН	560	-14.589 -52.793 168.443 1.00 49.18 S O	
30	MOTA	8316	0	НОН	561	14.274 -25.186 115.196 1.00 68.97 S O	
	ATOM	8317	0	НОН	562	-11.876 -66.758 99.635 1.00 58.70 S O	
	MOTA	8318	0	НОН	563	9.652 -71.240 90.805 1.00 47.55 S O	
	MOTA	8319	0	нон	564	8.337 -90.428 100.502 1.00 40.50 S O	
0.5	MOTA	8320	0	НОН	565	-4.356 -37.312 154.893 1.00 45.44 S O	
35	MOTA	8321	0	НОН	566	-11.188 -96.445 143.149 1.00 46.91 S O	
	MOTA	8322	0	нон	567	-15.389 -34.354 115.276 1.00 38.19 S O	
	MOTA	8323	0	нон	568	-6.686 -38.757 165.227 1.00 40.62 S O	
	MOTA	8324	0	нон	569	0.603 -92.714 105.614 1.00 49.80 S O	
40	MOTA	8325	0	нон	570	-10.908 -44.492 149.043 1.00 55.75 S O	
40	MOTA	8326	0	нон	571	11.715 -56.940 144.398 1.00 50.46 S O	
	MOTA	8327	0	HOH	575	14.024 -39.506 110.121 1.00 77.11 S O	
	MOTA	8328	0	нон	576	6.524 -55.615 104.387 1.00 50.00 S O	
	MOTA	8329	0	нон	577	9.972 -97.480 121.036 1.00 48.12 S O	
	MOTA	8330	0	HOH	578	-20.801 -42.139 108.101 1.00 41.31 S O	
45	MOTA	8331	0	HOH	580	-14.408 -77.621 170.420 1.00 47.84 S O	
	MOTA	8332	0	HOH	583	18.062 -48.141 147.872 1.00 46.19 S O	
	MOTA	8333	0	HOH	584	-11.118 -38.154 93.126 1.00 52.49 S O	
	MOTA	8334	0	HOH	585	18.723 -67.355 118.436 1.00 43.61 S O	
	MOTA	8335	0	HOH	586	18.807 -22.870 147.950 1.00 49.22 S O	
50	MOTA	8336	0	HOH	587	0.369 -10.834 151.620 1.00 48.37 S O	
	MOTA	8337	0	нон	590	-12.421 -41.425 151.484 1.00 54.81 S O	
	MOTA	8338		HOH	591	-6.884 -66.074 136.375 1.00 39.42 S O	
	MOTA	8339	0	HOH	592	0.492 -44.413 93.739 1.00 55.03 S C	
	ATOM	8340		нон	593	6.207 -11.926 150.116 1.00 51.39 S O	
55	ATOM	8341		нон	594	-15.591 -54.380 154.053 1.00 44.25 S C	
	ATOM	8342		HOH	595	11.989 -62.147 169.236 1.00 53.46 S C	)
	MOTA	8343	0	нон	597		)
	MOTA	8344	0	HOH	598	-4.024 -96.765 138.695 1.00 54.76 S C	)

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	ATOM	8345	0	нон	599	4.947	-13.288	156.213	1.00 41.25	s	0
	ATOM	8346	Ö	нон	601	-10.913			1.00 50.93	s	ŏ
	MOTA	8347	Ö	нон	603		-24.982		1.00 47.50	s	ō
	ATOM	8348	ŏ	нон	604		-94.049		1.00 42.28	s	ō
5	ATOM	8349	ŏ	нон	605	-14.289			1.00 52.11	s	ŏ
•	ATOM	8350	ŏ	нон	606	-19.924		99.672	1.00 57.18	s	ō
	MOTA	8351	ō	нон	607			123.118	1.00 46.97	š	ō
	ATOM	8352	ŏ	нон	608			115.087	1.00 40.46	s	ŏ
	MOTA	8353	ŏ	нон	609			107.395	1.00 55.14	s	ŏ
10	ATOM	8354	ō	нон	610			116.432	1.00 37.22	s	ō
	MOTA	8355	ŏ	нон	611			173.180	1.00 44.74	s	ō
	ATOM	8356	ŏ	нон	612			133.659	1.00 45.83	s	ŏ
	ATOM	8357	Ö	НОН	614			162.828	1.00 34.32	S	Ō
	ATOM	8358	ŏ	НОН	617			112.739	1.00 50.67	s	ō
15	ATOM	8359	ō	нон	618			169.501	1.00 48.06	s	ō
	ATOM	8360	ŏ	нон	619			148.628	1.00 54.49	s	ŏ
	ATOM	8361	ŏ	нон	620			136.844	1.00 53.29	s	ō
	ATOM	8362	ō	нон	621			164.880	1.00 46.75	s	ō
	ATOM	8363	ō	нон	623			148.740	1.00 58.12	s	ō
20	ATOM	8364	ō	нон	626			146.410	1.00 52.87	s	Ö
	MOTA	8365	ŏ	НОН	627			128.206	1.00 61.98	s	Ō
	ATOM	8366	ō	НОН	628			170.673	1.00 41.69	s	ō
	ATOM	8367	ō	НОН	629			155.302	1.00 39.69	S	Ō
	ATOM	8368	ŏ	НОН	630			116.783	1.00 48.41	s	ō
25	ATOM	8369	Ō	НОН	631			165.191	1.00 46.58	s	0
	ATOM	8370	Ō	НОН	633			169.718	1.00 41.32	S	Ō
	MOTA	8371	ō	НОН	634			111.912	1.00 48.12	S	Ō
	MOTA	8372	Ō	нон	635		-57.565		1.00 32.16	S	0
	MOTA	8373	ō	нон	636			147.829	1.00 42.76	S	Ō
30	ATOM	8374	Ō	нон	637			122.015	1.00 37.58	S	0
	ATOM	8375	0	нон	638			152.992	1.00 55.36	S	0
	MOTA	8376	0	нон	639			108.941	1.00 40.46	S	0
	MOTA	8377	0	нон	640	-12.826	-31.755	98.733	1.00 55.93	S	0
	MOTA	8378	0	HOH	641	-12.041	-90.235	104.096	1.00 48.46	S	0
35	ATOM	8379	0	HOH	642	-22.335	-75.465	150.732	1.00 50.86	S	0
	MOTA	8380	0	HOH	643	14.088	-18.781	123.781	1.00 49.25	S	0
	ATOM	8381	0	нон	644	-8.248	-95.373	103.761	1.00 53.33	S	0
	MOTA	8382	0	HOH	645	-8.607	-32.918	166.992	1.00 59.33	S	0
	MOTA	8383	0	HOH	646	5.063	-70.803	169.143	1.00 50.88	S	0
40	MOTA	8384	0	HOH	647	16.334	-45.651	141.775	1.00 62.45	S	0
	MOTA	8385	0	нон	648	-5.014	-44.649	141.840	1.00 53.76	S	0
	MOTA	8386	0	нон	649	-13.704	-26.229	101.833	1.00 51.97	S	0
	ATOM	8387	0	HOH	651			3 147.662	1.00 44.65	S	0
	MOTA	8388	0	HOH	653	-9.912	-68.068	3 134.022	1.00 53.05	S	0
45	MOTA	8389	0	HOH	655			3 138.891	1.00 54.56	S	0
	MOTA	8390	0	HOH	656			5 121.208	1.00 50.84	S	0
	MOTA	8391	0	HOH	658			l 144.738	1.00 47.88	S	0
	MOTA	8392	0	нон	659			1 110.573	1.00 41.41	S	0
<u></u>	MOTA	8393	0	HOH	661			1 102.774	1.00 47.84	S	0
50	MOTA	8394	0	HOH	663		-75.06		1.00 52.90	S	0
	MOTA	8395	0	HOH	664			7 101.333	1.00 57.41	S	0
	MOTA	8396		нон	668			3 170.230		S	0
	MOTA	8397		HOH	672			0 160.931		S	0
	MOTA	8398		нон	673			1 106.971		S	0
55	MOTA	8399		нон	674			3 106.975		S	0
	MOTA	8400		нон	676			1 110.887		S	0
	MOTA	8401		нон	677			1 104.475		S	0
	MOTA	8402	0	HOH	681	-10.652	2 -35.07	5 160.241	1.00 52.13	S	0

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	MOTA	8403	0	нон	685	-0.033	-82.942	139.985	1.00 54.94	s	0
	ATOM	8404	ŏ	нон		-12.713			1.00 70.33		Ō
	ATOM	8405	ŏ	нон		-10.374			1.00 55.51	s	0
	MOTA	8406	ŏ	нон		-15.464			1.00 50.64	s	Õ
5	MOTA	8407	ŏ	нон	691		-60.842		1.00 68.98	S	0
J	ATOM	8408	Ö	нон	692		-44.563		1.00 52.71	s	ō
	ATOM	8409	ŏ	нон	693		-94.906		1.00 42.37	S	0
	ATOM	8410	ŏ	нон	695		-62.997		1.00 43.89	S	0
	ATOM	8411	ŏ	нон	697		-62.955		1.00 67.72	s	Ö
10	MOTA	8412	Ö	нон	698		-84.834	96.026	1.00 68.94	s	O
••	ATOM	8413	ŏ	нон	701	-19.964			1.00 70.97	S	O
	ATOM	8414	ŏ	нон	702	-17.043			1.00 52.11	S	Ō
	ATOM	8415	ō	нон	703		-29.599		1.00 44.42	S	0
	ATOM	8416	ŏ	нон	708	-13.381		97.616	1.00 54.25	S	0
15	ATOM	8417	ŏ	нОн	709	-17.495			1.00 55.23	S	0
10	ATOM	8418	ŏ	нон	714		-18.552		1.00 43.63	s	Ö
	MOTA	8419	ŏ	НОН	719			126.958	1.00 51.95	S	0
	ATOM	8420	ŏ	нон	723			117.416	1.00 57.09	S	0
	MOTA	8421	Ö	нон	725	-20.095			1.00 48.48	S	Ö
20	MOTA	8422	ŏ	нон	728	-20.097		98.029	1.00 45.93	s	Ō
20	ATOM	8423	ő	нон	730			172.241	1.00 52.85	s	ō
	MOTA	8424	ő	нон	733		-77.427	94.470	1.00 48.88	S	Ō
	ATOM	8425	ŏ	нон	734			134.367	1.00 72.16	s	ō
	ATOM	8426	ŏ	нон	735			129.231	1.00 30.02	s	Ō
25	ATOM	8427	Ö	нон	736			125.987	1.00 26.22	S	0
20	MOTA	8428	ŏ	нон	737			100.638	1.00 41.15	s	Ō
	ATOM	8429	ő	нон	738			160.742	1.00 32.42	S	0
	ATOM	8430	ŏ	нон	739		-59.000	93.865	1.00 32.91	s	Ö
	ATOM	8431	Ö	нон	741			150.140	1.00 25.10	s	O
30	ATOM	8432	Ö	нон	742			155.807	1.00 40.46	s	Ō
00	MOTA	8433	Ö	нон	743			105.014	1.00 40.07	s	Ō
	ATOM	8434	ő	нон	744			105.086	1.00 29.75	S	Ō
	MOTA	8435	ŏ	НОН	745			144.159	1.00 34.27	S	0
	ATOM	8436	ŏ	нон	746			105.752	1.00 34.90	S	0
35	ATOM	8437	ŏ	нон	747			162.206	1.00 31.38	S	Ó
00	ATOM	8438	ŏ	нон	748			107.364	1.00 43.45	S	0
	ATOM	8439	Ö	нон	749		-39.637		1.00 60.23	S	0
	ATOM	8440	ō	нон	750			157.176	1.00 35.25	S	0
	ATOM	8441	Ö	нон	751			130.416	1.00 51.18	S	0
40	ATOM	8442	ŏ	нон	756			135.708	1.00 70.13	S	0
	ATOM	8443	Ö	нон	760			154.565	1.00 51.07	S	0
	ATOM	8444	_	нон	768			123.786	1.00 58.62	S	0
	ATOM	8445		нон	770			135.470	1.00 48.19	S	0
	ATOM	8446		нон	771			168.175	1.00 49.12	S	0
45	MOTA	8447		НОН	773			105.507	1.00 49.76	s	0
	MOTA	8448		нон	774			101.201	1.00 56.64	S	0
	MOTA	8449		нон	775			115.724	1.00 55.57	S	0
	ATOM	8450		нон	776		-49.262		1.00 33.00	S	0
	ATOM	8451		нон	777			160.148	1.00 54.59	S	0
50	ATOM	8452		нон	778			160.150	1.00 47.65	s	0
	ATOM	8453		нон	779	17.290	-20.49	5 151.229	1.00 42.84	S	0
	MOTA	8454		нон	783			5 106.538		S	0
	ATOM	8455		нон	784			125.976		S	0
	MOTA	8456		нон	785			3 129.815		s	0
55	ATOM	8457			786			7 129.222		s	0
-	ATOM	8458			787			0 134.871		s	0
	ATOM	8459			788			8 159.140			
	ATOM	8460			789			3 158.430			
			_								

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	ATOM	8461	0	нон	790	-15.720	-76 210	153 508	1.00	36 09	s	0
	ATOM	8462	ŏ	нон	791		-32.487		1.00		s	ŏ
	ATOM	8463	ŏ	нон	795	-15.122			1.00		S	o
	ATOM	8464	ŏ	нон	802		-43.210		1.00		S	o
5	ATOM	8465	Ö	нон	804		-52.044	92.549	1.00		S	Ö
9	ATOM	8466	0	НОН	805		-36.887		1.00		S	0
	ATOM	8467	0	НОН	806			140.997	1.00		S	Ö
	ATOM	8468	0	НОН	807	-14.220			1.00		S	Ö
	ATOM	8469	Ö	НОН	808			157.359	1.00		S	o
10	TER	8470	O	НОН	808	-10.210	-34.370	137.339	1.00	30.07	\$	0
10	ATOM	8471	C1	596	1	-2 439	_45 929	106.741	1.00	10 61	L	С
	ATOM	8472	N2	596	1			105.931	1.00			N
	ATOM	8473	C3	596	î			107.663	1.00		L.	C
	MOTA	8474	C4	596	1			106.859		19.46	L	C
15	MOTA	8475	C5	596	1			106.355		20.57	L	C
13	ATOM	8476	C6	596	1			100.333		18.69	L	C
	ATOM	8477	N7	596	1			104.338		19.90	L	N
	ATOM	8478	C8	596	1			108.636		18.27	L	C
	ATOM	8479	C9	596	1			107.869		19.94		C
20	ATOM			596	1			107.863			L	
20		8480		596	1			105.817		22.09	L	N
	MOTA	8481		596				103.486		18.66	L	C
	ATOM	8482			1			108.767		19.06	L	C
	MOTA	8483	C13	596 596	1					20.54	L	C
25	MOTA	8484		596 596	1			106.410		23.52	L	C
25	ATOM	8485			1			106.666		18.98	L	C
	MOTA	8486		596	1			107.482		20.47	L	N
	MOTA	8487		596	1			108.680		21.54	Ţ	0
	MOTA	8488		596	1			106.319		23.09	L	C
30	MOTA	8489		596	1			105.908		23.60	L	C
30	ATOM	8490		596	1			107.977		19.73	L	C
	ATOM	8491		596	1			106.476		18.94	r	C
	ATOM	8492		596	1			107.159		23.10	L	C
	MOTA	8493		596	1			105.437		23.56	L	C
25	MOTA	8494		596	1		-42.897			23.72	L	C
35	MOTA	8495		596	1			106.433		23.90	L	C
	ATOM	8496		596	1			109.074		20.03	L	C
	ATOM	8497		596	1			107.563		19.94	L	C
	MOTA	8498		596	1			107.082		23.17	L	C
40	ATOM	8499		596	1			105.363		23.29	L	C
40	ATOM	8500	C30		1			104.418		24.28	L	C
	ATOM	8501 8502		596 596	1			105.906		24.30 19.58	L	C
	ATOM	-			1			108.860			L	C
	MOTA	8503		596	1			106.184		23.09	L	C
45	MOTA	8504		596	1			104.890		23.94	L	C
45	MOTA	8505		596	1			106.157		19.63	L	Н
	ATOM	8506		596	1			104.677		19.04	L	H
	ATOM	8507		596	1			104.055		19.09	L	Н
	ATOM	8508		596	1		-46.507			18.86	L	Н
50	ATOM	8509		596	1		-42.349			22.39	L	Н
50	MOTA	8510		596	1			104.699		18.82	L	Н
	ATOM	8511		596	1			105.768		18.85	L	Н
	ATOM	8512		596	1			109.542		19.54	L	Н
	MOTA	8513		596	1			107.467		23.34	L	Н
EE	ATOM	8514		1 596	1			107.008		20.75	L	Н
55	ATOM	8515		5 596	1			107.553		20.71	L	Н
	ATOM	8516		5 596	1			5 108.126		19.63	L	H
	ATOM	8517		7 596	1			105.478		19.30	L	
	MOTA	8518	H4	3 596	1	-2.277	-40.85	7 107.889	1.00	23.21	L	Н

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	ATOM	8519	H49	596	1	0.389	-39.503	104.794	1.00 23.37	L	Н
	ATOM	8520	H50	596	1	0.781	-43.548	104.470	1.00 23.80	L	Н
	ATOM	8521	H51	596	1	2.062	-40.480	107.234	1.00 23.82	L	Н
	MOTA	8522	H52	596	1	-5.160	-44.208	110.074	1.00 19.98	L	Н
5	MOTA	8523	н53	596	1	-8.005	-45.966	107.398	1.00 19.79	L	н
	MOTA	8524	н54		1		-38.756		1.00 23.18	L	Н
	ATOM	8525	н55		1		-37.365		1.00 23.37	L	Н
	ATOM	8526	н56		1		-43.752		1.00 24.22	L	Н
	ATOM	8527	н57		î		-40.656		1.00 24.15	L	Н
10	ATOM	8528	н58		1		-45.633		1.00 19.69	L	Н
.0	ATOM	8529	н59		î		-36.980		1.00 13.05	L	H
	MOTA	8530	н60		1		-42.244		1.00 23.40	L	Н
	MOTA	8531	C1	596	2		-69.170		1.00 24.40	R	C
		8532	N2	596	2		-70.327		1.00 17.81		N
15	ATOM									R	
15	ATOM	8533	C3	596	2		-69.331		1.00 18.16	R	C
	ATOM	8534	C4	596	2			107.183	1.00 17.87	R	C
	MOTA	8535	C5	596	2			106.552	1.00 17.68	R	C
	MOTA	8536	C6	596	2			105.341	1.00 16.75	R	С
~~	MOTA	8537	N7	596	2			107.475	1.00 18.02	R	N
20	MOTA	8538	C8	596	2			108.702	1.00 17.84	R	C
	MOTA	8539	C9	596	2	•		108.148	1.00 18.61	R	С
	MOTA	8540		596	2			106.004	1.00 18.38	R	N
	MOTA	8541		596	2			105.985	1.00 16.15	R	С
22	MOTA	8542		596	2			108.904	1.00 17.98	R	С
25	ATOM	8543	C13	596	2	5.614	-65.865	108.504	1.00 18.62	R	C
	ATOM	8544	C14	596	2	2.047	-73.304	106.474	1.00 18.34	R	С
	ATOM	8545	C15	596	2	7.255	-71.129	107.206	1.00 16.15	R	С
	ATOM	8546	N16	596	2	6.925	-65.908	108.153	1.00 18.90	R	N
	ATOM	8547	017	596	2	5.159	-64.930	109.159	1.00 20.09	R	0
30	MOTA	8548	C18	596	2	2.661	-74.677	106.388	1.00 17.44	R	C
	MOTA	8549	C19	596	2	0.683	-73.086	105.780	1.00 20.03	R	С
	ATOM	8550	C20	596	2	6.750	-71.375	108.506	1.00 16.48	R	С
	ATOM	8551	C21	596	2	8.371	-70.304	107.048	1.00 15.06	R	C
	ATOM	8552	C22	596	2	3.682	-74.969	107.305	1.00 16.86	R	С
35	ATOM	8553	C23	596	2	2.335	-75.659	105.424	1.00 17.31	R	С
	ATOM	8554	C24	596	2	0.509	-72.121	104.758	1.00 19.86	R	С
	ATOM	8555		596	2			106.157	1.00 21.14	R	С
	ATOM	8556	C26	596	2			109.617	1.00 16.65	R	C
	ATOM	8557		596	2			108.142	1.00 15.10	R	C
40	ATOM	8558		596	2			107.242	1.00 16.83	R	Č
	ATOM	8559		596	2			105.375	1.00 17.45	R	č
	ATOM	8560		596	2			104.108	1.00 21.12	R	Č
	ATOM	8561		596	2			105.487	1.00 22.11	R	C
	MOTA	8562		596	2			109.421	1.00 16.40	R	C
45	ATOM	8563		596	2			106.291	1.00 17.12	R	
40	MOTA	8564		596	2			104.454	1.00 17.12		
		8565		596	2			104.434	1.00 21.79	R R	C
	MOTA				2						Н
	MOTA	8566		596				105.110	1.00 16.89	R	Н
EΩ	ATOM	8567		596	2			104.403	1.00 16.85	R	Н
50	MOTA	8568		596	2		-68.493		1.00 18.19	R	Н
	MOTA	8569		596	2			105.274	1.00 18.51	R	
	MOTA	8570		596	2			105.252	1.00 16.39	R	
	MOTA	8571		596	2			106.242	1.00 16.29	R	
	ATOM	8572		596	2			109.657	1.00 18.45	R	
55	MOTA	8573		596	2			107.522	1.00 18.65	R	
	MOTA	8574		596	2			107.652		R	
	MOTA	8575		596	2			108.408		R	
	MOTA	8576	H46	596	2	5.903	-72.034	108.633	1.00 16.43	R	H

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	MOTA	8577	H47	596	2		-70.133		1.00 15.62	R	H
	MOTA	8578	H48	596	2	3.925	-74.260	108.081	1.00 16.88	R	Н
	MOTA	8579	H49	596	2	1.556	-75.498	104.697	1.00 17.51	R	H
	MOTA	8580	H50	596	2	1.310	-71.479	104.438	1.00 20.27	R	Н
5	MOTA	8581	H51	596	2	-0.390	-74.586	106.951	1.00 21.25	R	Н
	MOTA	8582	H52	596	2	6.891	-70.827	110.605	1.00 16.51	R	Н
	MOTA	8583	н53	596	2	9.797	-69.006	108.010	1.00 15.29	R	Н
	MOTA	8584	H54		2		-76.405		1.00 17.12	R	Н
	ATOM	8585		596	2		-77.634		1.00 17.47	R	Н
10	ATOM	8586	н56		2		-71.205		1.00 21.25	R	Н
. •	ATOM	8587	H57		2		-74.336		1.00 22.07	R	н
	ATOM	8588	н58		2		-69.329		1.00 16.27	R	н
	ATOM-	8589		596	2		-78.095		1.00 10.27	R	Н
	ATOM	8590	H60	596	2		-72.657		1.00 17.47	R	Н
15											
15	MOTA	8591	C1	596	3		-46.116		1.00 15.32	T	C
	MOTA	8592	N2	596	3		-44.983		1.00 16.26	T	N
	ATOM	8593	C3	596	3		-45.890		1.00 15.57	T	С
	MOTA	8594	C4	596	3			157.515	1.00 14.61	$\mathbf{T}$	C
	MOTA	8595	C5	596	3		-44.165		1.00 16.95	T	С
20	MOTA	8596	C6	596	3			159.405	1.00 15.77	$\mathbf{T}$	C
	MOTA	8597	N7	596	3			156.986	1.00 14.76	T	N
	MOTA	8598	C8	596	3	1.501	-46.839	155.702	1.00 14.30	$\mathbf{T}$	C
	ATOM	8599	C9	596	3	3.340	-48.270	156.482	1.00 15.09	$\mathbf{T}$	C
	MOTA	8600	N10	596	3	1.720	-42.818	158.524	1.00 16.85	${f T}$	N
25	MOTA	8601	C11	596	3	5.028	-43.656	158.829	1.00 15.65	$\mathbf{T}$	С
	MOTA	8602	C12	596	3	2.274	-48.030	155.565	1.00 14.20	$\mathbf{T}$	C
	ATOM	8603	C13	596	3	4.091	-49.540	156.364	1.00 13.71	$\mathbf{T}$	C
	MOTA	8604		596	3			157.922	1.00 16.81	$\mathbf{T}$	С
	ATOM	8605		596	3			157.706	1.00 16.86	T	Ċ
30	ATOM	8606		596	3			156.931	1.00 12.82	$\bar{\mathbf{T}}$	N
•	ATOM	8607		596	3			155.803	1.00 12.64	T	Ö
	ATOM	8608	C18		3			157.929	1.00 15.58	T	Č
	ATOM	8609		596	3			158.482	1.00 16.81	T	c
	ATOM	8610		596	3			156.366	1.00 17.32	Ť	c
35	ATOM	8611		596	3			157.971	1.00 17.32	T	C
33	MOTA	8612		596	3			156.970	1.00 17.30	T	C
				596	3			158.869		Т	C
	ATOM	8613			3				1.00 15.97 1.00 17.21		
	ATOM	8614		596				159.499		T	C
40	MOTA	8615		596	3			157.976	1.00 17.34	T	C
40	MOTA	8616		596	3			155.318	1.00 17.94	T	C
	MOTA	8617		596	3			156.911	1.00 18.26	T	C
	ATOM	8618		596	3			156.961	1.00 15.61	T	С
	MOTA	8619		596	3			158.836	1.00 16.06	T	С
4.5	ATOM	8620		596	3			160.051	1.00 17.06	T	С
45	MOTA	8621		596	3			158.511	1.00 17.58	${f T}$	C
	MOTA	8622		596	3			155.595	1.00 17.92	T	C
	MOTA	8623		596	3			157.872	1.00 15.78	$\mathbf{T}$	С
	MOTA	8624		596	3			159.552	1.00 17.41	Т	C
	MOTA	8625	н35	596	3	4.396	-47.438	158.229	1.00 14.89	$\mathbf{T}$	H
50	MOTA	8626	н36	596	3	4.548	-45.560	159.719	1.00 16.02	T	H
	MOTA	8627	н37	596	3	3.570	-44.278	3 160.303	1.00 16.08	T	Н
	ATOM	8628		596	3			154.998	1.00 14.65	Т	н
	ATOM	8629		596	3			3 159.303	1.00 17.20	T	Н
	ATOM	8630		596	3			159.607	1.00 16.13	T	Н
55	ATOM	8631		596	3			7 158.496	1.00 16.08	T	H
	ATOM	8632		596	3			154.791	1.00 14.79	T	Н
	ATOM	8633		596	3			2 156.879	1.00 16.80	T	Н
	ATOM	8634		596	3			157.389	1.00 10.60	T	
	AIOM	0034	uaa	. 230	3	J. / I	,	. 131.303	1.00 12.05	T.	Н

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	MOTA	8635	H45 596	3	5.897	-50.359	156.902	1.00 12.70	т	Н
	MOTA	8636	H46 596	3		-43.403		1.00 17.56	T	Н
	MOTA	8637	H47 596	3		-45.259		1.00 17.47	T	Н
	ATOM	8638	H48 596	3		-41.118		1.00 15.78	T	Н
5	ATOM	8639	H49 596	3		-39.633		1.00 16.06	T	Н
	ATOM	8640	н50 596	3		-43.510		1.00 17.16	T	Н
	MOTA	8641	H51 596	3		-40.340		1.00 17.30	T	Н
	ATOM	8642	H52 596	3		-44.629		1.00 17.79	T	Н
	MOTA	8643	H53 596	3		-46.371		1.00 17.79	T	Н
10	MOTA	8644	H54 596	3		-39.108		1.00 15.13	T	Н
. •	ATOM	8645	H55 596	3		-37.532		1.00 15.83	T	Н
	ATOM	8646	H56 596	3		-43.416		1.00 15.79	T	
	MOTA	8647	H57 596	3		-40.295				Н
	ATOM	8648	H58 596	3		-46.063		1.00 17.59	T	Н
15	MOTA	8649	H59 596					1.00 18.12	T	Н
13	ATOM	8650		3		-37.276		1.00 15.95	T	Н
	ATOM	8651		3		-41.800		1.00 17.61	T	H
				4		-68.974		1.00 15.41	V	C
	MOTA	8652	N2 596	4		-70.123		1.00 15.55	V	N
20	MOTA	8653	C3 596	4		-69.217		1.00 16.29	V	С
20	ATOM	8654	C4 596	4		-67.768		1.00 15.23	V	С
	ATOM	8655	C5 596	4		-70.970		1.00 17.80	V	С
	ATOM	8656	C6 596	4		-70.373		1.00 14.78	V	С
	MOTA	8657	N7 596	4			156.854	1.00 17.03	V	N
05	MOTA	8658	C8 596	4		-68.243		1.00 15.66	V	С
25	MOTA	8659	C9 596	4			156.248	1.00 16.16	V	С
	MOTA	8660	N10 596	4			158.283	1.00 19.21	V	N
	MOTA	8661	C11 596	4			158.336	1.00 15.05	V	С
	MOTA	8662	C12 596	4			155.502	1.00 15.69	V	С
20	MOTA	8663	C13 596	4	-5.921	-65.520	155.976	1.00 15.40	V	С
30	MOTA	8664	C14 596	4			157.784	1.00 20.54	V	C
	MOTA	8665	C15 596	4			157.105	1.00 14.31	V	С
	MOTA	8666	N16 596	4			156.246	1.00 15.48	V	N
	MOTA	8667	017 596	4			155.454	1.00 16.46	V	0
25	MOTA	8668	C18 596	4			157.842	1.00 20.15	V	С
35	MOTA	8669	C19 596	4			158.435	1.00 21.79	V	С
	ATOM	8670	C20 596	4			155.847	1.00 14.50	V	С
	MOTA	8671	C21 596	4			157.201	1.00 13.56	V	C
	MOTA	8672	C22 596	4			156.878	1.00 19.47	V	C
40	MOTA	8673	C23 596	4			158.809	1.00 20.11	V	С
40	ATOM	8674	C24 596	4			159.483	1.00 22.05	V	С
	ATOM	8675	C25 596	4			157.954	1.00 22.52	V	С
	MOTA	8676		4			154.701	1.00 15.23	V	С
	ATOM	8677	C27 596	4			156.046	1.00 14.11	V	С
45	MOTA	8678	C28 596	4			156.897	1.00 20.11	V	С
45	MOTA	8679	C29 596	4			158.792	1.00 19.91	V	С
	MOTA	8680	C30 596	4			160.033	1.00 22.72	V	С
	MOTA	8681	C31 596	4			158.513	1.00 23.89	V	С
	MOTA	8682	C32 596	4			154.798	1.00 14.57	V	С
50	ATOM	8683	C33 596	4			157.857	1.00 19.08	V	С
50	MOTA	8684	C34 596	4			159.561	1.00 23.31	V	С
	MOTA	8685	н35 596	4	-6.566	-67.610	157.776	1.00 15.39	V	H
	MOTA	8686	н36 596	4	-6.869	-69.448	159.250	1.00 15.16	V	Н
	MOTA	8687	н37 596	4			159.890	1.00 15.23	v	Н
	MOTA	8688	н38 596	4			155.105	1.00 15.91	V	Н
55	MOTA	8689	н39 596	4			158.987	1.00 19.56	V	Н
	ATOM	8690	H40 596	4			159.057	1.00 14.81	V	H
	ATOM	8691	H41 596	4	-6.976	-72.235	158.063	1.00 14.91	V	Н
	MOTA	8692	H42 596	4	-3.664	-66.262	154.828	1.00 16.36	V	Н

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	MOTA	8693	H43	596	4	-3.023	-73.063	156.736	1.00	20.49	V	Н
	MOTA	8694	H44	596	4	-7.674	-66.265	156.678	1.00	15.81	V	H
	MOTA	8695	H45	596	4	-7.746	-64.660	156.003	1.00	15.79	V	H
	ATOM	8696	H46	596	4	-6.745	-71.667	155.781	1.00	14.88	V	Н
5	ATOM	8697	H47	596	4	-9.601	-69.601	158.167	1.00	14.04	V	Н
	MOTA	8698	H48	596	4	-5.019	-74.086	156.124	1.00	19.70	V	Н
	ATOM	8699	H49	596	4	-2.932	-75.479	159.612	1.00	20.34	V	H
	ATOM	8700	H50	596	4	-2.242	-71.633	159.911	1.00	22.23	V	Н
	ATOM	8701	H51	596	4	-0.914	-74.727	157.153	1.00	22.61	V	Н
10	ATOM	8702	H52	596	4	-7.560	-70.482	153.747	1.00	15.36	V	Н
	ATOM	8703	н53	596	4	-10.538	-68.503	156.125	1.00	14.26	V	Н
	MOTA	8704	H54	596	4	-6.380	-76.155	156.163	1.00	19.84	V	Н
	ATOM	8705	н55	596	4	-4.177	-77.575	159.533	1.00	20.19	V	Н
	ATOM	8706	H56	596	4	-0.042	-71.464	160.825	1.00	23.01	V	Н
15	ATOM	8707	H57	596	4	1.346	-74.536	158.135	1.00	23.57	V	H
	ATOM	8708	н58	596	4	-9.501	-68.923	153.917	1.00	14.82	V	H
	ATOM	8709	н59	596	4	-5.939	-77.923	157.887	1.00	19.84	V	Н
	λπOM	9710	460	596	Δ	1 775	-72 930	160.006	1.00	23.48	v	Н

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## References

The references listed below as well as all references cited in the specification are incorporated herein by reference to the extent that they supplement, explain, provide a background for or teach methodology, techniques and/or compositions employed herein.

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It will be understood that various details of the invention can be changed without departing from the scope of the invention. Furthermore, the foregoing description is for the purpose of illustration only, and not for the purpose of limitation, the invention being defined by the claims.